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- (54) BLOCK COPOLYMER
- (57) Provided are a block copolymer including a polymer block containing acrylonitrile or methacrylonitrile as a principal constituent, which is excellent in heat resistance, weatherability, oil resistance, flame retardancy, and low-temperature resistance and which can be economically produced; and a thermoplastic resin com-

position and an elastomer composition each containing the block copolymer. The block copolymer is produced by reversible addition-fragmentation chain transfer polymerization in the presence of a thiocarbonylthio groupcontaining compound.

Description

Technical Field

[0001] The present invention relates to block copolymers. More particularly, the invention relates to a block copolymer including a polymer block which contains a monomer selected from the group consisting of acrylonitrile and methacry-lonitrile as a principal constituent. Furthermore, the present invention relates to a thermoplastic resin composition or elastomer composition containing the block copolymer as an essential component.

10 Background Art

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[0002] Materials in which acrylonitrile or methacrylonitrile is polymerized have been widely used because of their excellent heat resistance, weatherability, oil resistance, flame retardancy, etc. Commonly known examples thereof include acrylonitrile-styrene-acrylic rubber copolymer resins (AAS), acrylonitrile-ethyrene styrene copolymer resins (ASS), acrylonitrile-butadiene-styrene copolymer resins (ABS), acrylonitrile-butadiene-styrene copolymer resins (ASS), acrylonitrile-butadiene-copolymer rubber (NBR), and hydrogenated attitle nubber.

[8003] However, all of these materials are formed by random copolymertzation of acrylonitrile and other monomers or by graft copolymerization of acrylonitrile unto polymer chains. With respect to block copolymers including polymer blocks containing acrylonitrile and methacrylonitrile as principal constituents, although research has been conducted on the laboratory level, no production has been performed industrially.

[0004] In general, in an acrylonitrile-containing polymer, as the acrylonitrile content is increased, oil resistance, was resistance, heat resistance, heat resistance, each strength are improved, while low-temperature resistance tends to be degraded. In contrast, in a block copolymer in which a polymer block containing acrylonitrile or methacrylonitrile as a principal constituent and a flexible polymer block composed of other constituents, low-temperature resistance is considered to be improved while maintaining excelence in least resistance, of resistance, etc.

[0005] As the methods for synthesizing block copolymers, living polymerization methods are usually used. Examples of living polymerization methods include living cationic polymerization methods, living anionic polymerization methods, and living radical polymerization methods. Among them, living radical polymerization methods are most useful because of the applicability to a wide variety of monomers and also because of the applicability to water-based polymerization. Commonly known examples of the living radical polymerization methods include a method using nitroxyl radicals, such as 2.2.6.6-tetramethyl-1-piperidinyloxy (TEMPO) radicals, described in Junpo He et al., Polymer, 2000, 41, p.4573; and an atom transfer radical polymerization method described in Matyjaszewski et al., J. Am. Chem. Suc., 1995, 117, p.5614. However, in the method using nitroxyl radicals, little has been reported about the polymerization of monomers other than styrene, and no example has been known about the polymerization of acrylonitrile. Furthermore, in general, since the polymerization temperature must be 120°C or more, this method is not economical for industrial production. On the other hand, in the atom transfer radical polymerization, since nitrile groups coordinate to a metal complex which is a catalyst, polymerization does not proceed satisfactorily. Furthermore, a complicated purification step is required in order to remove the metal complex from the polymer, thus being uneconomical. Additionally, in the living radical polymerization methods described above, it is generally difficult to perform water-based polymerization, such as emulsion polymerization or suspension polymerization. In the living radical polymerization methods described above, there are cases in which it is difficult to perform water-based polymerization, such as emulsion polymerization or suspension polymerization.

45 Disclosure of Invention

[0006] The present invention has been achieved to overcome the problems associated with the conventional techniques. It is an object of the present invention to provide a block copolymer including a polymer block containing acylonitile or methacylonitile as a principal constituent, which is excellent in heat resistance, weatherability, oil resistance, flame retardancy, and low-temperature resistance and which can be economically produced. It is another object of the present invention to provide a thermoplastic resin composition or elastomer composition containing the block copplymer.

[0007] A block copolymer of the present invention includes a polymer block (A) and a polymer block (B), the block copolymer being produced by forming the polymer block (A) by reversible addition-fragmentation chain transfer polymerization in the presence of a thiocarbonylthio group-containing compound, and then by forming the polymer block

[9008] The polymer block (A) is prepared by (co)polymerizing 50% to 100% by weight of at least one monomer selected from the group consisting of acrylonitrile and methacrylonitrile and 50% to 0% by weight of at least one mon-

omer selected from the group consisting of methacrylate esters, styrene, and α-methylstyrene.

[0009] The polymer block (B) is prepared by (co)polymerizing at least one monomer selected from the group consisting of acrylic acid, methacrylic acid, acrylate esters, methacrylate esters, vinyl acetate, styrene, α -methylstyrene, butadiene, isoprene, and vinyl chloride.

1,0010] A block copolymer of the present invention may be produced by separately forming the polymer block (A) and the polymer block (B) each by a reversible addition-fragmentation chain transfer polymerization method in the presence of a thiccarbonyithio group-containing compound, and then by coupling the polymer block (A) and the polymer block (B). 10011] The block copolymer of the present invention can be of various types. Examples of types include, but are not limited to, an A-B diblock copolymer including the polymer block (A) and the polymer block (B), an A-B-A thiblock copolymer. a R-B-B thiblock copolymer, and a branched block acceptioner. Polymers produced by linking monomers other than the ones described above to such polymers may also be acceptable. Examples thereof include an A-B-C polymer including a polymer chain C formed by a monomer other than the ones described above.

[0012] The block copolymer of the present invention is produced by a reversible addition-fragmentation chain transfer (RAFT) method in the presence of a thiocarbonythic group-containing compound. RAFT polymerization methods a disclosed, for example, in PCT Publication No. WO98/01478; PCT Publication No. WO99/01478; PCT Publication No. WO99/01414; Macromolecules, 1998, 31, p.5559; Macromolecules, 1999, 32, p.2071; Macromolecules, 1999, 32, p. 977; and Macromolecules, 1999, 32, p. 977. and 1997. and 1

1013] That is, the monomers for forming the polymer block (A) are polymerized in the presence of a thiocarbonylthio group-containing compound to form the polymer block (A). The monomers for forming the polymer block (B) inited to the polymer block (A). Thereby, a block copolymer including the polymer block (A) and the polymer block (B) is produced. In this method, the thiocarbonylthio group-containing compound functions as a chain transfer agent, and the resultant block copolymer contains a tleast one thiocarbonylthio group in each molecule. The block copolymers including the polymer blocks (A) and the polymer blocks (B) are a multiblock copolymer, such as the A-B-A or B-A-B type, or a multiblock copolymer, such as the A-B-A or B-A-B type, or a multiblock copolymer, such as the A-B-A or B-A-B type, or a formal polymer block (B) are a copoling and then by coupling the polymer block (A) and the polymer block (B) separately, each in the presence of a thiocarbonylthio group-containing compound, and then by coupling the polymer block (A) and the polymer block (A) and the polymer block (B) are the p

[0014] When the block copolymers, or the polymer blocks (A) and (B), are coupled to each other, the block copolymers preferably have functional groups at the ends in view of the fact that coupling can be performed easily and reliably. In the present invention, the thiocarbonythio groups introduced into the polymer by the method described above can be used as such functional groups.

[0015] When the thiocarbonylthio group-containing copolymers are subjected to a coupling reaction, for example, first, the copolymers are allowed to react with a processing agent composed of at least one compound selected from the group consisting of bases, acids, and hydrogen-hitrogen bond-containing compounds to convert the thiocarbonylthio groups of the copolymers into mercapto groups or mercaptide groups. Next, coupling of the copolymers is performed using the mercapto groups or mercaptide groups. The block copolymer with the desired type is thereby obtained.

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[0016] The polymer block (A) is prepared by (co)polymerizing 50% to 100% by weight of at least one monomer selected from the group consisting of acrylonitrile and methacytonitrile and 50% to 0% by weight of at least one monomer selected from the group consisting of methacytale esters, styrene, and c-methylstyrene.

[0017] Because of the excelience in oil resistance and flame retardancy of the resultant block copolymer, the polymer block (A) is prepared preferably by polymerizing 80% to 100% of at least one monomer selected from the group consisting of acryoinstrille and methacrylonitrile and 20% to 0% of at least one monomer selected from the group consisting of methacrylate esters, styrene, and c-methylstyrene, and more preferably by polymerizing 100% by weight of at least one monomer selected from the group consisting of acrylonitrile and methacrylonitrile. With respect to the amounts of acrylonitrile and methacrylonitrile used, in view of availability and cost, preferably 50% by weight or more of acrylonitrile is polymerized, more preferably 80% by weight or more of acrylonitrile is polymerized, and most preferably 100% by weight of archylonitrile is polymerized.

[0018] In view of the strength and workability of the resultant block copolymer, the molecular weight distribution, which is determined by get permeation chromatography analysis, of the polymer block (A) is preferably 1.8 or tess, and more preferably 1.6 or tess, and more preferably 1.6 or tess, in the present invention, the molecular weight distribution (Mw/Mn) is a value corresponding to the ratio of the weight-average molecular weight (Mw) to the number-average molecular weight (Mn) determined by get permeation chromatography analysis.

55 [0019] In view of the strength and workability of the resultant block copolymer; the number-average molecular weight of the polymer block (A) is preferably in the range of 1,000 to 500,000, and more preferably in the range of 3,000 to 100,000.

[8820] Because of the excellence in heat resistance and strength of the resultant block copolymer, the glass transition

temperature of the polymer block (A) is preferably 50°C or more, and more preferably 80°C or more. In the present invention, the glass transition temperature (Tg) is determined according to the following equation:

$$(1/Tg) = (W_2/Tg_1) + (W_2/Tg_2) + ... + (W_m/Tg_m)$$

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(wherein Tg_1 , Tg_2 , ..., Tg_m are the glass transition temperatures correspond to monomers used for polymerization when they are homopolymerized, respectively, W_1 , W_2 , ..., W_m are the weight fractions of the monomer components used for polymerization in the polymer block, respectively, W_1 and the relationship $W_1 + W_2 + ..., W_m = 1$ is satisfied). Additionally, the above equation is generally known as the Fox equation. With respect to the glass transition temperatures of the individual homopolymens (Tg_1 , Tg_2 , ..., Tg_m), for example, the values described in Polymer Handbook Third Edition (Wiley-Interscience, 1989) may be used.

[0021] Among the monomers constituting the polymer block (A), as at least one monomer selected from the group consisting of methacrylate esters, styrene, and a-methylstyrene, because of the excellence in oil resistance, weatherability, and heat resistance, methocrylate esters are preferable. Examples of such methocrylate esters include, but are not limited to, alkyl esters, anyl esters, and aralkyl esters of methacrylic acid. Examples of methacrylate esters include, but are not limited to, methyl methacrylate, ethyl methacrylate, n-butyl methacrylate, isobutyl methacrylate, tert-butyl methacrylate, 2-ethylhexyl methacrylate, tridecyl methacrylate, stearyl methacrylate, cyclohexyl methacrylate, benzyl methacrylate. 2-hydroxyethyl methacrylate, 2-hydroxypropyl methacrylate, diethylaminoethyl methacrylate, glycldyl methacrylate, tetrahydrofurfurvi methacrylate, ethylene glycol dimethacrylate, triethylene glycol dimethacrylate, letraethylene glycoi dimethacrylate, 1,3-butylene glycol dimethacrylate, trimethylolpropane trimethacrylate, isopropyl methacrylate, pentyl methacrylate, hexyl methacrylate, heptyl methacrylate, octyl methacrylate, nonyl methacrylate, decyl methacrylate, dodecyl methacrylate, phenyl methacrylate, tolyl methacrylate, isobornyl methacrylate, 2-methoxyethyl methacrylate, 3-methoxybutyl methacrylate, 2-aminoethyl methacrylate, 2-methacryloyloxypropyltrimethoxysilane, 2-methacryloyloxypropyldimethoxymethylsilane, trifluoromethyl methacrylate, pentafluoroethyl methacrylate, 2-aminoethyl methacrylate, 2-(N,N-dimethylamino)ethyl methacrylate, 3-aminopropyl methacrylate, 3-(N,N-diethylamino) propyl methacrylate, and 2,2,2-trifluoroethyl methacrylate. These compounds may be used alone or in combination. Among these methacrylate esters, because of the excellence in heat resistance, weatherability, oil resistance, and strength of the resultant copolymers, methacrylate esters of alcohols containing 4 or less carbon atoms, such as methyl methacrylate, ethyl methacrylate, and n-butyl methacrylate, are preferable, and methyl methacrylate is particularly preferable.

[0022] Additionally, by adding methacrylate esters having functional groups, when the resultant block copolymers are molded, crosslinking reactions can be carried out using the functional groups. Thereby, molded objects having excellent strength and compression set can be produced. It is also possible to produce molded objects having excellent compatibility with various resins and rubbers using the functional groups. Examples of such functional groups include, but are not limited to, a hydroxyl group, an amino group, an epoxy group, a carboxyl group, and a crosslinkable silyl group. Examples of the methacrylate esters containing such functional groups include, but are not limited to, hydroxyl group-containing methacrylate esters, such as 2-hydroxyethyl methacrylate and 2-hydroxypropyl methacrylate, amino group-containing methacrylate esters, such as 2-aminoethyl methacrylate and 3-aminopropyl methacrylate; epoxy group-containing methacrylate esters, such as glycidyl methacrylate; carboxyl group-containing methacrylate esters. such as 3-methacryloyloxypropanoic acid and 4-methacryloyloxybenzoic acid; and crosslinkable silyl group-containing methacrylate esters, such as 2-methacryloyloxypropyltrimethoxysilane and 2-methacryloyloxypropyldimethoxymethyisilane. Alternatively, after the monomers are polymerized, carboxyl groups may be introduced by hydrolysis. These methacrylate esters may be used alone or in combination. If the content of the functional group-containing methacrylate ester is too high, it may be brittle when the block copolymer is formed and crosslinked. Therefore, the content of the functional group-containing methacrylate ester is preferably less than 10% by weight, and more preferably less than 5% by weight, of the polymer block (A). However, this does not apply to the case when functional groups are introduced in order to improve compatibility with various resins, rubbers, etc.

[0023] As described above, the polymer block (B) in the block copolymer of the present invention is composed of at least one monomer selected from the group consisting of acrylic acid, methacrylic acid, acrylate esters, methacrylate esters, winy acetate, styrene, a-methylstyrene, butadlene, isoprene, and vnnyl chloride. These monomers may be used alone or in combination. Additionally, after the polymerization, the vinyl acetate unit may be saponified, the butadlene unit or isoprene unit may be hydrogeneted, or the butadlene unit, sorpene unit, or vinyl chloride unit may be chlorinated. [0024] Among the monomers constituting the polymer block (B), examples of acrylate esters include, but are not limited to, alkyl esters, anyl esters, and aralkyl esters of acrylic acid. Examples of acrylate esters include, but are not imited to, methyl acrylate, ethyl acrylate, h-propyl acrylate, isopropyl acrylate, h-butyl acrylate, hexyl acrylate, 2-thyl-hexyl acrylate, boxponyl acrylate, 2-thyloxyethyl acrylate, boxponyl acrylate, 2-thydroxyptha crylate, boxponyl acrylate, 2-thydroxyptha crylate, 2-thydroxyptha crylate, 2-thydroxyptha acrylate, 3-thydroxyptha crylate, 2-thydroxyptha acrylate, 3-thydroxyptha crylate, 3-thydroxyptha crylate, 2-thydroxyptha crylate, 3-thydroxyptha crylate, 3-thydroxy

pyl acylate, stearyl acylate, glyoddy acylate, 2-acyloyloxypropyldimethoxymethylsiaine, 2-acyloyloxypropyltrimethroxysilane, trifluoromethyl acylate, pentafluoroethyl acylate, 2-(2,2-trifluoroethyl acylate, 3-dimethylaminoethyl acylate, sobulyl acylate, 4-hydroxybulyl acylate, scrylate of alkyl-modified dipenteerylhritol, ethylene oxide-modified bisphenol A diacylate, carbitol acylate, acylate of e-caprolactone-modified dipenteerylhritol, ethylene actone-modified etteralydrotruryl acylate, diacylate of acprolactone-modified dipenteerylhritol, caprolactone-modified etteralydrotruryl acylate, diacylate of acprolactone-modified repentily glycol hydroxypiates, distinuethylolpropane ethracylate, diacylate, diacylate, branchylolpropane ethoralydrotruryla cylate, dipenteerylhritol hexacorylate, tinethylolpropane ethoralydrotruryla erylate, diacylate, diacrylate of neopentyl glycol hydroxypivatate, 1,9-nonandiol acylate, 1,4-butanediol acylate, 2-propanoic acid [2-1,1-dimethyl-2-(1-oxo-2-propenylloxy)glythyl-5-clyt-1,3-dioxane-5-yll methyl ester, 1,6-hexanediol acylate, acylate, 3-denalpathyliol tracylate, 2-denalpathylioloxypidyliolydrogen phthalate, methyl 3-methyl ester, 1,6-hexanediol acylate, 2-penalpathyliolitolydroxyliolydroxypidyliolydrogen phthalate, methyl 3-methyl ester, 1,6-hexanediol acylate, 2-denalpathyliolitolydroxyliolydroxypidyliolydroxyliolydr

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[9025] Among the monomers constituting the polymer block (B), the methacrylate ester is not particularly limited.

Examples of methacrylate esters are the same as those described above with reference to the polymer block (A).

[0026] Among the monomers constituting the polymer block (B), in view of heat resistance and weathereality, acrylate esters are preferable; in view of flexibility, acrylate sesters are preferable; and in view of availability and cost, acrylate esters are preferable. The view of flexibility acrylate, and tort-butyl acrylate, and tort-butyl acrylate are more preferable. When the block copolymers of the present invention are used as textiles, in view of texture, surface properties, and flame retardancy, vinyl choride is preferable beautiful and polycarbonate resins, preferred monomers for use are as follows: acrylic acid, methacrylic acid, functional group-containing acrylate esters, and functional group-containing acrylate esters, and supplementations are used as extensional propositions are preferable. The present invention are used as extensional propositions are preferable as the present invention are used as follows: acrylic acid, methacrylic acid, functional group-containing acrylate esters, such as hydroxyl group-containing acrylate esters, such as polydroxyl group-containing methacrylate esters, are propositionally acrylate esters, amino group-containing methacrylate esters, are propositionally acrylate este

[0027] The monomers constituting the polymer block (B) are preferably selected based on the following criteria. For example, because of the excellence in flexibility and low-temperature resistance of the resultant block copolymer, the glass transition temperature of the polymer prepared by polymertzing the monomers only is 30°C or less, and more preferably 0°C or less. The glass transition temperature of the polymer block (B) is determined according to the Fox

equation as in the polymer block (A).

[1028] In view of the strength and workability of the resultant block copolymer, the molecular weight distribution, which is determined by get permeation chromatography analysis, of the polymer prepared by polymerizing only the monomers constituting the polymer block (B) is preferably 1.8 or less, and more preferably 1.5 or less.

[0029] Furthermore, in view of flexibility and elasticity, the number-average molecular weight, which is determined by gel permeation chromatography analysis, of the polymer prepared by polymerizing only the monomers constituting the polymer block (B) is preferably in the range of 1,000 to 1,000,000, more preferably in the range of 3,000 to 500,000, and most preferably in the range of 5,000 to 200,000.

[0030] As described above, the block copolymer of the present invention is produced by forming the polymer block (A) and then by forming the polymer block (B), and furthermore by coupling them to each other. Alternatively, the polymer block (B) and the polymer block (B) are formed separately, and then coupling is performed. In view of workability and the excellence in compatibility when the block copolymer is mixed with resins, rubbers, or the like, the molecular welpht distribution, which is determined by eal permeasion chromatography analysis, of the block copolymer is preferably 1.8 or less, and more preferably 1.8 or less.

[0031] The thiocarbonylthio group-containing compound used as a chain transfer agent in the radical polymerization described above is at least one compound selected from the group consisting of a compound represented by general formula (1).

$$\left(\begin{array}{c}
S\\Z^1-C-S-R^1
\end{array}\right) \qquad (1)$$

(wherein R1 is a p-valent organic group of 1 or more carbon atoms which may contain one of hitrogen, oxygen, sulfur, halogen, silicon, phosphorus, and metal atoms, or which may be a polymer; Z1 is a hydrogen atom, halogen atom, monovalent organic group of 1 or more carbon atoms which may contain one of hitrogen, oxygen, sulfur, halogen, silicon, and phosphorus atoms, or which may be a polymer; when plural Z1s are present, the piural Z1s may be the same or different and p is an integer of 1 or more), and a compound represented by general formula (2):

(wherein R² is a monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, suffur, halogen, silicon, phosphorus, and metal atoms, or which may be a polymer; 2^2 is an oxygen atom (when q = 2), or $q = 2^2$, suffur atom (when q = 2), or $q = 2^2$, suffur atom (when q = 2), and $q = 2^2$, or $q = 2^2$, suffur atom (when q = 2), or $q = 2^2$, or $q = 2^2$, and $q = 2^2$, and q

[0032] In the compound having the thiocarbony(thio structure represented by general formula (1), R1 is not particularly limited. In view of availability of the compound, preferably, R1 has 1 to 20 carbon atoms, and p is 6 or less. Examples of R1 include alkyl, autibution alkyl, autibution are represented by polyalent aliphatic hydrocarbon group, a polyvalent aliphatic hydrocarbon group with an airphatic group, a polyvalent aliphatic hydrocarbon group with an airphatic group, and polyvalent aliphatic hydrocarbon group with an airphatic group, and a polyvalent aromatic with a profession and polyvalent aromatic substituted hydrocarbon group containing a heteroatom, and a polyvalent aromatic substituted hydrocarbon group containing a heteroatom. In view of availability of the compound and polymerization activity, R1 is preferably benzyl. 1-phenylethyl, 2-(2-phenylpropyl), 1-act-butyl, 1,1,3,3-tetramathyl-butyl, 2-(2-phorophenylpropyl), hydrocarbon group or profession group service and profession groups represented by the following formulae.

$$O$$
 $-CH_2-CO+CH_2CH_2O+CH_3$

$$-CH_2 - CO + CHCH_2O + CH_3$$

$$CN$$
 Q CH_3 $-C$ CH_2CH_2 CH_3 CH_3

 $\begin{array}{c} \text{CN} \\ -\text{C--}\text{CH}_2\text{CH}_2\text{COOH} \\ \text{CH}_3 \end{array}$

20 (In the formulae, n is an integer of 1 or more, and r is an integer of 9 or more.)

$$CH_2$$
 CH_2
 CH_2
 CH_2
 CH_2

--(CH₂)_n--

(In the formulae, R4 is a divatent organic group of 1 or more carbon atoms; n is an integer of 1 or more; r is an integer of 0 or more; and plural r's may be the same or different.)

(In the formulae, n is an integer of 1 or more; r is an integer of 0 or more; and plurel r's may be the same or different.) [0033] In the above formulae, in view of availability of the compound, preferably, each of n and r is 500 or less. Preferably, R4 has 1 to 20 carbon atoms. Examples of R4 Structures include, but are not limited to, -{CH₂}_n- (wherein n is an integer of 1 or more). C₈H₄-, and -CH₂-C₈H₄-CH₂-.

[0034] In the compound having the thiocarbonylithio structure represented by general formula (1), Z² is not particularly limited. In view of availability of the compound and polymerization activity, when Z⁴ is an organic group, preferably to organic group has 1 to 20 across nations. Examples of Z⁴ include alkyl, substituted alkyl, substituted arkyl, are substituted arkyl, are substituted arkyl, substituted arakyl, heterocyclic, N-aryl-N-alkylamino, N-N-dialylamino, NN-dialkylamino, thiobalkyl, and dialkylphosphinyl groups. In view of availability of the compound, Z⁴ is preferably phenyl, methyl, ethyl, benzyl, 4-dinorphenyl, 1-naphtyl, Z-arablythyl, diethosylhosphinyl, n-buyl, tert-butyl, methoxy, ethoxy, thiomethyl (methylsulfide), phenoxy, thiophenyl (phenylsulfide), NN-dimethylamino, N-N-diethylamino, N-phenyl-N-methylamino, N-phenyl-N-ethylamino, thiobenzyl (benzylsulfide), pentafluorophenoxy, or any one of organic groups represented by formulae below.

[0035] In the compound having the thiocarbony/thio structure represented by general formula (2), R² is not particularly limited. In view of availability of the compound and polymerization activity, preferably, R² has 1 to 20 carbon adrous Examples of R² include alkyl, substituted aralkyl, and substituted aralkyl, In view of availability of the compound, R² is preferably benzyl, 1-phenylethyl, 2-(2-phenylpropyl), 1-acetoxyethyl, 1-(4-methoxyphenylpethyl, ethoxycarbonylpethyl, 2-(2-cyanobropyl), includencyl, tert-butyl, 1.1,3-letramethy/butyl, 2-(2-(p-chlorophenylpropyl), vinylbenzyl, tert-butylsulfide, 2-carboxylethyl, carboxytmethyl, cyanomethyl, 1-cyanoethyl, 2-(2-cyanobutyl), or any one of organic groups represented by formulae below, (wherein n is an integer of 1 or more, and r is an integer of 0 or more).

$$\begin{array}{ccc} & CH_3 \\ -CH_2 - CO + CHCH_2O + CH_3 \end{array}$$

$$CN$$
 O CH_3 $-C$ $-CH_2CH_2-CO$ $CHCH_2O$ r CH_3

[0036] In the above formulae, each of n and r is preferably 500 or less in view of availability of the compound.

[0037] In the compound having the thiocarbonylthio structure represented by general formula (2), 2° is not particularly imited. In view of availability of the compound and polymerization activity, when Z° is an organic group, preferably, the organic group has 1 to 20 carbon atoms, and a is 6 or less. Examples of Z° include a polyvalent alighatic hydrocarbon group, a polyvalent aromatic hydrocarbon group, a polyvalent alighatic hydrocarbon group with an aromatic ing, a polyvalent aromatic hydrocarbon group with an eliphatic group, a polyvalent alighatic hydrocarbon group containing, a heteroatom, and a polyvalent aromatic substituted hydrocarbon group containing a heteroatom. In view of availability of the compound, preferably Z° has any one of the structures represented by formulae below, (wherein n is an integer of 1 or more, and r is an integer of 0 or more).

[9038] In the above formulae, each of n and r is preferably 500 or less in view of availability of the compound.
[9039] Among the thiocarbonylthio group-containing compounds used in the present invention, in view of availability and the fact that:
A B discount of the discount of the present invention.
A B discount of the present invention, in view of availability and the fact that:

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$$z^{1}$$
— C — S — R^{2} (3)

(wherein R² is a monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, phosphorus, and metal atoms, or which may be a polymer, and Z1 is a hydrogen atom, halogen atom, or monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, and phosphorus atoms, or which may be a polymer). In view of availability of the compound and polymerization activity, preferably, R² has 1 to 20 carbon atoms. The specific examples of the structures of R² and Z¹ in the formula are the same as those described with reference to R² in general formula (2) and Z¹ in general formula (1), respectively.

[0040] In the present invention, when the polymer blocks (A) and (B) are synthesized separately and then are coupled to each other to produce an A-B-A triblock copolymer, because of ease in production, preferably, a compound represented by general formula (B) is used as the thiocarbonythio group-containing compound used for the preparation of the polymer block (A); and a compound represented by general formula (4) below is used as the thiocarbonythio group-containing compound used for the preparation of the polymer block (B);

$$S$$
 S $Z^1-C-S-R^3-S-C-Z^1$ S

(wherein R³ is a divalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, hallogen, silicon, phosphorus, and metal atoms, or which may be a polymer; each Z¹ is a hydrogen atom, hallogen atom, or monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, and phosphorus atoms, or which may be a polymer; and two Z¹s may be the same or different). Specific examples of Z¹ structures in the formula are the same as those described with reference to Z¹ in general formula (1).

[041] In the thiocarbonylthio group-containing compound represented by general formula (4), preferably, R⁵ has 1 to 20 carbon atoms. Although R⁵ is not particularly limited, in view of availability and polymerization activity, preferably. R⁵ has any one of the structures represented by formulae below.

$$\begin{array}{c} Q & Q \\ -\text{CH}_2\text{CO}-(\text{CH}_2\text{CH}_2\text{O})_{\overline{\textbf{h}}}-\text{R}^4-(\text{OCH}_2\text{CH}_2)_{\overline{\textbf{h}}}-\text{OC}-\text{CH}_2-\text{CH}_2-\text{CH}_2)_{\overline{\textbf{h}}} \end{array}$$

(In the above formulae, R4 is a divalent organic group of 1 or more carbon atoms; n is an integer of 1 or more; r is an integer of 0 or more; and plural r's may be the same or different.)

[0042] In the above formulae, each of n and r is preferably 500 or less in view of availability of the compound. Preferably, R² has the 20 carbon atoms. Examples of R³ structures include, but are not limited to, $\neg (CH_2)_{n_1}$ (wherein is an integer of 1 or more), $\neg (AH_2)_{n_1}$ (wherein is an integer of 1 or more).

[0043] Specific examples of compounds having the thiocarbonylthio structures, which are used in the present invention, include, but are not limited to, compounds represented by formulae below, (wherein Me, Et, Ph, and Ac represent

methyl, ethyl, phenyl, and acetyl, respectively; R4 is a divalent organic group of 1 or more carbon atoms; n is an integer of 1 or more; r is an integer of 0 or more; and plural r's may be the same or different).

$$Ph-C-S-CH_2COO(CH_2CH_2O)Me$$

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[0044] In the above formulae, each of n and r is preferably 500 or less in view of availability of the compound. Preferably, R⁴ has 1 to 20 carbon atoms. Examples of R⁴ structures include, but are not limited to, -(CH₂)_n- (wherein is an integer of 1 or more). C-2H_x-, and -CH₂-C₈-H₃-CH₂-C

[0045] The amount of the thiocarbonythio group-containing compound used in the present invention is not particularly limited and can be stoichiometrically activated based on the monomers used. In general, since the number of motor of the resultant polymer is substantially equal to the number of moles of the thiocarbonythio group-containing compound, the molecular weight of the polymer can be controlled by adjusting the molar ratio of the monomers to the thiocarbonythiol group-containing compound. When the rate of reaction of the monomers is 100%, the theoretical molecular weight of the resultant polymer is represented by (x/y) × Min + Mir, wherein Min is the molecular weight of the monomers used, it is the number of moles of the monomers used, Mr is the molecular weight of the thiocarbonythiol group-containing compound used. Consequently, the amount of the thiocarbonythiol group-containing compound used.

[0046] The technique used for radical polymerization is not particularly limited when the block copolymer of the present invention is prepared. Any method commonly used in the art, such as bulk polymerization, solution polymerization, emulsion polymerization, suspension polymerization, or microsuspension polymerization, may be employed. Among them, in view of cost and safety, water-based polymerization, such as emulsion polymerization, as uspension polymerization, or microsuspension polymerization, is preferred.

[0047] In the case of solution polymerization of the monomers, examples of solvents which may be used include, but are not limited to, hydrocarbon solvents, such as heptane, hexane, octane, and mineral spirit; ester solvents, such as ethyl acetate, n-butyl acetate, isobutyl acetate, ethylene glycol monomethyl ether acetate, and diethylene glycol monobulyl ether acetate; ketone solvents, such as acetone, methyl ethyl ketone, methyl isobutyl ketone, dilsobutyl ketone, and cyclohexanone; alcohol solvents, such as methanol, ethanol, isopropanol, n-butanol, sec-butanol, and isobutanol; ether solvents, such as tetrahydrofuran, diethyl ether, dibutyl ether, dioxane, ethylene glycol dimethyl ether, and ethylene glycol diethyl ether; amide solvents, such as dimethylformamide, diethylformamide, dimethylacetamide, and diethylacetamide; and aromatic petroleum solvents, such as toluene, xylene, benzene, Swasol 310 (manufactured by Cosmo Oil Co., Ltd.), Swasol 1000 (manufactured by Cosmo Oil Co., Ltd.), and Swasol 1500 (manufactured by Cosmo Oil Co., Ltd.). These solvents may be used alone or in combination. The types and amounts of solvent used may be determined in consideration of the solubility of the monomers used, the solubility of the resultant polymer, the polymerization initiator concentration and the monomer concentration suitable for achieving a satisfactory reaction rate, the solubility of the thiocarbonylthic group-containing compound, effects on human body and environment, availability, cost, etc., and are not particularly limited. In view of solubility, availability, and cost, industrially, toluene, dimethylformamide, tetrahydrofuran, and acetone are preferable, and dimethylformamide and toluene are more preferable. [9048] In the present invention, in the case of emulsion polymerization or microsuspension polymerization of the monomers, examples of emulsifiers which may be used include, but are not limited to, anionic surfactants, such as fatty acid soap, rosin acid soap, sodium naphthalenesulfonate-formalin condensates, sodium alkylsulfonate (e.g., so-

dium dodecyl sulfonate), sodium alkylbenzene sulfonate, sodium alkylsulfate (e.g., sodium dodecyl sulfate), ammonium alkylsulfate, triethanolamine alkylsulfate, sodium dialkylsulfosuccinate, sodium alkyldiphenylether disulfonate, sodium polyoxyethylene alkyl ether sulfate, and sodium polyoxyethylene alkylphenyl ether sulfate; nonionic surfactants, such as polyoxyethylene alkyl ether, polyoxyethylene higher alcohol ether, sorbitan fatty acid ester, polyoxyethylene sorbitan faity acid ester, polyoxyethylene sorbitol fatty acid ester, glycerin fatty acid ester, polyoxyethylene fatty acid ester, polyoxyethylene aklylamine, and alkyl alkanolamide; and cationic surfactants, such as alkyltrimethylammonium chloride. These emulsifiers may be used alone or in combination. As necessary, a cationic surfactant, such as an alkylamine hydrochloride, may be used, or a dispersant for suspension polymerization which will be described below may also be added. The amount of the emulsifier used is not particularly limited. Because of good emulsification state and the fact that polymerization proceeds smoothly, the amount of the emulsifier is preferably 0.1 to 20 parts by weight based on 100 parts by weight of the monomers. Among the emulsifiers described above, in view of stability of the emulsification state, nonionic surfactants are preferable. Additionally, in order to stabilize the emulsification state, various emulsifying aids may also be used. Examples of emulsifying aids include, but are not limited to, linear hydrocarbons, such as hexadecane; hydrocarbon polymers, such as polyethylene, polypropylene, polybutadiene, and hydrogenated polybutadiene: polar grganic solvents, such as acetone, ethanol, and methanol; and higher alcohols, such as octyl alcohol, decyl alcohol, and lauryl alcohol. The amount of the emulsifying aid used is not particularly limited. Because of excellence in the balance between cost and effect, the amount of the emulsifying aid used is preferably 0.1 to 20 parts by weight, and more preferably 0.5 to 15 parts by weight, based on 100 parts by weight of the emulsifier.

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[8049] In the case of suspension polymerization of the monomers, any dispersant commonly used may be used Examples of dispersants include, but are not limited to, pertailly seponified polytiving leastenty, polytiving alcohol), methyl cellulose, carboxymethyl cellulose, gelatin, poly(alfytiene oxide), and combinations of anionic surfactants and dispersing agents. These may be used alone or in combination. The emulsifier used for emulsion polymerization described above may also be used as necessary. The amount of the dispersant used is not particularly limited. Because of the fact that polymerization proceeds smoothly, the amount of the dispersant is preferably 0.1 to 20 parts by weight based on 100 parts by weight of the monomers used.

[0050] The polymerization initiator or polymerization initiation method used for the radical polymerization is not particularly limited, and any polymerization initiator or polymerization initiation method commonly used may be employed. Examples of polymerization initiators include, but are not limited to, peroxide polymerization initiators, such as methyl ethyl ketone peroxide, methyl isobutyl ketone peroxide, cyclohexanone peroxide, methyl cyclohexanone peroxide, isobutyryl peroxide, 3.5.5-trimethylhexanovi peroxide, laurovi peroxide, benzovi peroxide, tert-butyl hydroperoxide, cumene hydroperoxide, diisopropylbenzene hydroperoxide, p-menthane hydroperoxide, 1,1,3,3-tetramethylbutyl hydroperoxide, di-tert-butyl peroxide, tert-butyl-α-cumyl peroxide, di-α-cumyl peroxide, 1,4-bis[(tert-butylperoxy)lsopropyljbenzene, 1,3-bis[(tert-butylperoxy)isopropyljbenzene, 2,5-dimethyl-2,5-bis(tert-butylperoxy)hexane, 2,5-dimethyl-2.5-bis(tert-butylperoxy)-3-hexyne, 1.1-bis(tert-butylperoxy)-3,3,5-trimethylcyclohexane, n-butyl-4,4-bis(tert-butylperoxy)valerate, 2,2-bis(tert-buty/peroxy)butane, tert-buty/peroxy acetate, tert-buty/peroxy isobuty/ate, tert-buty/peroxy octoale, terl-butylperoxy pivalate, tert-butylperoxy neodecanoate, tert-butylperoxy-3,5,5-trimethyl hexanoate, tertbutylperoxy benzoate, tert-butylperoxy laurate, 2,5-dimethyl-2,5-bis(benzoylperoxy)hexane, bis(2-ethylhexyl)peroxy dicarbonate, disopropylperoxy dicarbonate, di-sec-butylperoxy dicarbonate, di-n-propylperoxy dicarbonate, bis (3-methoxybutyl)peroxy dicarbonate, bis(2-ethoxyethyl)peroxy dicarbonate, bis(4-tert-butylcyclohexyl)peroxy dicarbonate, O-tert-butyl-O-isopropylperoxy carbonate, and succinic acid peroxide; azo polymerization initiators, such as 2.2'azobis-(2-amidinopropane)dihydrochloride, dimethyl 2,2'-azobis(Isobutyrate), 2,2'-azobis-(4-methoxy-2,4-dimethylvaleronitrile), 2,2'-azobis(isobutyronitrile), 1,1'-azobis(cyclohexane-1-carbonitrile), azocumene, 2,2'-azobis(2-methylbutyronitrile), 2,2'-azobis-(2,4-dimethylvaleronitrile, 4,4'-azobis(4-cyanovaleric acid), 2-(tert-butylazo)-2-cyanopropane, 2,2'-azobis(2,4,4-trimethylpentane), and 2,2'-azobis(2-methylpropane); inorganic peroxides, such as potassium persulfate and sodium persulfate; vinyl monomers which thermally generate radical species, such as styrene; compounds which generate radical species by light, such as benzoin derivatives, benzophenone, acylphosphine oxide, and photoredox systems; and redox polymerization initiators including sodium sulfite, sodium thiosulfate, sodium formaldehyde sulfoxylate, ascorbic acid, ferrous sulfate, or the like, as a reducing agent, and potassium peroxydisulfate, hydrogen peroxide, tert-butyl hydroperoxide, or the like, as an oxidizing agent. These polymerization initiators may be used alone or in combination. If may also be possible to use a polymerization initiation system by electron irradiation, X-ray irradiation, radiation irradiation, or the like. With respect to polymerization initiation methods using such initiators, the methods described in Moad and Solomon "The Chemistry of Free Radical Polymerization". Pergamon, London, 1995, pp. 53-95 may be employed.

[0051] In the present invention, the amount of the polymerization initiator used is not particularly limited. In order to produce a polymer with a narrow molecular weight distribution, the amount of radical species generated during polymerization is preferably 1 mole or less, and more preferably 0.5 moles or less, relative to 1 mole of thiocarbonyithio group in the thiocarbonyithio group-containing compound, in order to control the amount of radical species generated during polymerization, in addition to the control of the amount of the polymerization initiator, preferably, temperature is con-

trolled in the case of the polymerization initiator which causes thermal dissociation, or the amount of energy is controlled in the case of the polymerization initiation system which generates radicals by light or electron beams. Because of ease of control of polymerization, using a polymerization initiator which causes thermal dissociation, the polymerization reaction is carried out preferably at temperatures which allow the polymerization initiator to have a half-life of 0.5 to 50 hours, more preferably at temperatures which allow the polymerization initiator to have a half-life of 1 to 20 hours, and most preferably at temperatures which allow the polymerization initiator to have a half-life of 5 to 15 hours.

[0052] The block copolymer of the present invention is prepared using the solvent, emulsifier, polymerization initiator, etc., as will be described below.

[0053] The thiocarbonythio group-containing block copolymer is treated with a processing agent, as required, so that the thiocarbonythio groups are converted into mercapto groups or mercaptide groups. The processing agent used in the process is not particularly limited, in view of high yield, preferably, a method is employed in which the thiocarbonythio group-containing compound is allowed to react with a processing agent composed of a compound selected from the group consisting of bases, acids, and hydrogen-nitrogen bond-containing compounds. Among him, when a base or an acid is used, in the presence of water, thiocarbonythio groups are converted into mercapto groups by hydrolysis. When a hydrogen-nitrogen bond-containing compound is used, the presence of water is not required, which is preferable.

[0054] Examples of bases which may be used as processing agents include, but are not limited to, alkali metal hydroxides, such as sodium hydroxide, potassium hydroxide, and lithium hydroxides, are calcium hydroxides, and casium hydroxides, and advantage and a control of the control of the casium hydroxides, and a sodium methylate, sodium ethylate, and tilhium butylate, and altihium butylate; alkaline-ath metal altoxholates, such as smagnesium ethylate and magnesium ethylate and magnesium ethylate, and aluminum borohydride; and organometalin reagents, such as hydroxides, such as magnesium aluminum hydride, and aluminum borohydride; and organometalin reagents, such as hydroxidite, n-butylithium, tert-butylithium, ethylmagnesium bromide, and phenylmagnesium bromide. Furthermore, alkali metals, such as metallic indiction, and metallic potassium; and alkaline-earth metals, such as metallic magnesium and metallic calcium may also be used inome appeared to the control of the control of

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lithium hydroxide, calcium hydroxide, magnesium hydroxide, sodium methylate, and sodium ethylate. (10055) Examples of acids which may be used as processing spents include, but are not limited to, inorganic acids, such as hydrochloric acid, hitrò acid, suffuric acid, phosphoric acid, hydrofluoric acid, hydrofluoric acid, fluoroboric acid, chlorosulfonic acid, sindro acid, acid, acid, and silicofluoric acid, organic acids, such as p-foluenesulfonic acid, tiffuoromethy sulfonic acid, acid,

6 [0055] Examples of hydrogen-nitrogen bond-containing compounds which may be used as processing agents include, but are not limited to, ammonia, hydrazine, primary amine compounds, secondary amine compounds, amine hydrochlorides, hydrogen-nitrogen bond-containing polymers, and hindered amine light stabilizers (HALSs).

[0037] Among the hydrogen-nitrogen bond-containing compounds, specific examples of primary amine compounds include, but are not limited to, N-(2-aminoethy)ethanolamine, 12-aminododecanole acid, 3-amino-1-propanol, allylamine, isopropylamine, 3,3-ininobis(propylamine), monoethylamine, 2-ethylhexylamine, 3-(2-ethylhexyloxy)propylamine, 3-ethylamine, 3-(diethylamino)propylamine, 3-ethylamine, in-butylamine, in-butylamine, sec-butylamine, n-propylamine, 3-(methylamino)propylamine, 3-(diethylamino)propylamine, 1-ethylamine, N-methyl-3,3-iminobis(propylamine), 3-methoxypropylamine, 2-eminoethanol, ethylenedamine, diethylenehexioryloneyamine, N-carboxypropylamine, 1-eth-aminobutane, 1,2-diaminopropane, 1,3-diaminopropane, diaminomaleonitrile, cyclohexylamine, ATU (manufactured by Ajinomoto Co., Inc.), itilourea dioxide, 2-hydroxyethylaminopropylamine, hexamethylenedamine, n-hexamethylind-aramine, monomethylamine, mono

pylamine, antisidne, antiine, p-aminoacetanilide, p-aminobenzoic add, ethyl p-aminobenzoate ester, 2-amino4-chorophenol, 2-aminothizaole, 2-aminothiophenol, 2-aminothiophenol, 2-aminothiophenol, 2-aminothiophenol, 2-aminothiophenol, 2-aminothiophenol, 2-aminothiophenol, 2-aminothiophenol, 2-aminothiophenol, 2-dischiophenologica, 4-amino-5-hydroxy-2,7-naphthalenesulfonic acid, 3-dischiophenologica, 4-amino-6-hydroxy-2-paphthalenesulfonic acid, 3-dischiophenologica, 4-diaminositionic acid, 2-amino-5-aphthol-7-suffonic acid, 4-diaminonic p-cresidine, 6-diamino-3,3-diathyldiphenylmethane, 4.4-diaminobenzanilide, 6-diaminodiphenyl ether, 3,3-dimethyl-4,4-diaminodiphenylmethane, sulfanilic acid, tobias acid, 2,4-5-trichioroaniline, 0-tolidine, tolujdine, toluylenediamine, sodium naphthionate, antitroaniline, m-nitro-p-toluidine, 0-chloro-p-toluidine, m-sulfonic acid, phenythydrazine, phenylenediamine, phenetidine, phenethylamine, benzylamine, ben

[9058] Among the hydrogen-nitrogen bond-containing compounds, specific examples of secondary amine compounds include, but are not fimiled to, N-methylethanolamine, dislaylamine, dislayoropylamine, diethylamine, d

[0059] Among the hydrogen-nitrogen bond-containing compounds, specific examples of amide compounds include but are not limited to, 2-acrylamido-2-methylpropanesulfonic acid, dihydrazide adipate, N-isopropylacrylamide, N-t-octylacrylamide, carbohydrazides, guanythiourea, glycylglycne, N-l3-(dimethylamino)propyllamide, N-N-methylenebis (stearoamide), amide cleate, amide stearate, N-N-methylenebis (stearoamide), h-N-pyrozymethylstearoamide, diacetone ecrylemide, thioacetoamide, thioacethoydrazide, thiosemi-carbazide, thiourea, dihydrazide addecanedioate, dihydrazide adipate, dihydrazide sebacate, dihydrazide isophthalate, 1,5-hexamethylenebis(N-N-dimethylsemicarbazide), formamide, methacrylamide, N-N-methylenebis(acrylamide), N-methylolacrylamide, acetalide, acet

[0860] Among the hydrogen-nitrogen bond-containing compounds, specific examples of amine hydrochlorides inoldude, but are not limited to, acetamidine hydrochloride, 2.2-'azobis-(2-amidinopropane) dihydrochloride, monomethylamine hydrochloride, dimethylamine hydrochloride, monoethylamine hydrochloride, diethylamine hydrochloride, monopropylamine hydrochloride, dipropylamine hydrochloride, aminoguanidine hydrochloride, dibutylamine hydrochloride, semicarbazide hydrochloride, gualadine hydrochloride, aminoguanidine hydrochloride, 2-chloroethylamine hydrochloride, cysteamine hydrochloride, and ter-butyl hydrazine monohydrochloride.

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46 [0061] Among the hydrogen-nitrogen bond-containing compounds, specific examples of hydrogen-nitrogen bond-containing polymers include, but are not limited to, POLYMENT (manufactured by Nippon Shokubai Co., Ltd.), poly (ethylene inine), amino poly(acryl amide), nylon 6, nylon 66, nylon 610, nylon 612, nylon 11, nylon 12, nylon MXD6, nylon 48, polyamide-imide, polyandamine, and polyurethane.

[0062] Among the hydrogen-nitrogen bond-containing compounds, examples of HALSs include, but are not limited to. Adekasutabu LA-77 (manufactured by Asahi Denka Co., Ltd.), Chimassorb 944LD (manufactured by Ciba Specialty Chemicals), Adekasutabu LA-57 (manufactured by Ciba Specialty Chemicals), Adekasutabu LA-57 (manufactured by Asahi Denka Co., Ltd.), Adekasutabu LA-68 (manufactured by Asahi Denka Co., Ltd.), Adekasutabu LA-68 (manufactured by Asahi Denka Co., Ltd.), Adekasutabu LA-68 (manufactured by Asahi Denka Co., Ltd.), and Goodrite UV-3034 (manuf

[0063] When the thiocarbonyithio group-containing polymer is treated with the processing agent, the amount of the processing agent used is not particularly limited. When an aciditic compound or basic compound is used as the processing agent, in view of ease of handling and reactivity, the amount used is preferably 0.01 to 100 parts by weight, more preferably 0.05 to 50 parts by weight, and most preferably 0.1 to 30 parts by weight and most preferably 0.1 to 30 parts by weight by weight by weight preferably 0.05 to 50 parts by weight and most preferably 0.1 to 30 parts by weight by weight preferably 0.05 to 50 parts by weight parts preferably 0.05 to 50 parts by weight parts preferably 0.05 to 50 parts by weight preferably 0.05 to 50 parts by we

the polymer. When the polymer is treated with an acidic compound or basic compound, in view of stability of the polymer, preferably neutralization is carried out after treatment. When a hydrogen-nitrogen bond-containing compound is used as the processing agent, because of a high introduction rate of mercapto groups, the amount of the hydrogen-nitrogen bond-containing compound is preferably 0.5 to 1,000 motes, and more preferably 1 to 500 moles, based on 1 mole of hilocarbonythio group in the polymer. The excess hydrogen-nitrogen bond-containing compound can be recovered and reused. It is also possible to remove the excess processing agent by passing it through a column packed with an absorbent, such as silica or alturina.

[0064] Among the processing agents described above, preferred are hydrogen-nitrogen bond-containing compounds in view of no corrosion of apparatus, the fact that neutralization is not required, and stability of crosslinkable sityl groups. More preferred are ammonia, primary amine compounds with a boiling point of 100°C or less, and secondary amine compounds with a boiling point of 100°C or less in view of the fact that the purification step after treatment can be simplified. Most preferred are ammonia, monomethylamine, dimethylamine, monoethylamine, and diethylamine in view of availability.

[0055] In the present invention, when the thiocarbonylithic group-containing vinyl polymer is treated with the processing agent, the reaction conditions are not particularly limited. For example, a method in which the polymer is dissolved in an organic solvent, and the processing agent is added thereto; a method in which the processing agent is added to a water-based dispersion or emulsion; or a method in which the processing agent is directly added to the solid or molten polymer itself may be employed. The treatment temperature is not particularly limited. In view of reactivity, the treatment temperature is preferably -50°C to 300°C, and more preferably -10°C to 200°C.

[9066] By using the processing agent, the block copolymers of the present invention, which have mercapte groups or mercaptide groups, are obtained.

[0087] The block copplymers having mercapto groups or mercaptide groups at ends are formed by treatment with the processing agent. If necessary, the block copplymers thus obtained are subjected to a coupling reaction to produce a block copplymer of the desired type. For exemple, A-B diblock copplymers, each including a polymer block A and a polymer block B linked to the block A, are prepared, and by coupling the A-B diblock copplymers, block copplymers, such as A-B-A-b block copplymers and (A-B) block copplymers, are prepared.

[9068] Examples of coupling methods used for coupling the block copolymers having mercapto groups or mercaptide groups include, but are not limited to, methods (i) to (xi) for block copolymers having mercapto groups, and methods (xii) to (xv) for block copolymers having mercaptide groups. Examples of coupling methods for block copolymers having mercapto groups include (i) a method in which disulfide bonds are formed between block copolymers in the presence of an oxidizing agent, such as oxygen, lead dioxide, or calcium dioxide, and thereby the block copolymers are coupled; (ii) a method in which a compound having at least two isocyanato groups in each molecule is allowed to react with block copolymers, and thereby the block copolymers are coupled via thiourethane bonds (-NHCOS-); (iii) a method in which a compound having at least two isothlocyanato groups in each molecule is allowed to react with block copolymers. and thereby the block copolymers are coupled via dithiourethane bonds (-NHCSS-); (iv) a method in which a compound having at least two unsaturated bonds in each molecule is added to block copolymers, and thereby the block copolymers are coupled; (v) a method in which dehydrocondensation is performed between a polyvalent carboxylic acid and block copolymers, and thereby the block copolymers are coupled via thioester bonds; (vi) a method in which transesterification is performed between a polyvalent carboxylate ester and block copolymers, and thereby the block copolymers are coupled via thicester bonds: (vII) a method in which esterification is performed between a polyvalent carboxylic anhydride and block copolymers, and thereby the block copolymers are coupled via thioester bonds; (viii) a method in which acylation is performed between a polyvalent acyl halide and block copolymers, and thereby the block copolymers are coupled via thioester bonds; (ix) a method in which block copolymers are coupled by transesterification between a carbonate compound and the block copolymers; (x) a method in which a ketone is allowed to react with block copolymers to form thickets) bonds, and thereby the block copolymers are coupled; and (xi) a method in which block copolymers are coupled by dehydrocondensation between a compound having at least two hydroxyl groups in each molecule and the block copolymers. Examples of coupling methods for block copolymers having mercaptide groups include (xli) a method in which a compound having at least two halogen atoms in each molecule is allowed to react with block copolymers to form sulfide bonds (Williamson reaction), and thereby the block copolymers are coupled; (xiiil) a method in which block copolymers are coupled by neutralization between a polyvatent carboxylic acid and the block copolymers; (xiv) a method in which block copolymers are coupled by a reaction between a polyvalent carboxylic acid halide and the block copolymers; and (xv) a method in which disulfide bonds are formed between block copolymers in the presence of an oxidizing agent, and thereby the block copolymers are coupled.

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[0069] In addition, it is also possible to use compounds having different functional groups in each molecule. Examplies of such compounds include, but are not limited to, ketones having an isocyanato group in each molecule, compounds having an isocyanato group and an alkenyl group in each molecule, compounds having an isocyanato group and a halogen atom in each molecule, compounds having an example isocyanato group and an alikentic isocyanato group in each molecule, compounds having an aromatic isocyanato group and an aliphatic isocyanato group in each molecule, compounds having

a carboxyl group and a halogen atom in each molecule, compounds having a hydroxyl group and a halogen atom in each molecule, compounds having a mercapto group and a halogen atom in each molecule, compounds having a halogen atom and an alkenyl group in each molecule, compounds having an alkenyl group and a mercapto group in each molecule, and compounds having an isocyanato group and an ethinyl group in each molecule.

[9070] Among these methods, in view of ease of reaction and coupling efficiency, methods (i), (iii), (xii), and (xv) are preferred.

[9971] Among the coupling methods for block copolymers having mercapto groups, when method (i), in which disulfide bonds are formed between block copolymers in the presence of an oxidizing agent, is employed, examples of oxidizing agents which may be used include, but are not limited to, chlorates, such as sodium chlorate, potassium chlorate, ammonium chlorate, barium chlorate, and calcium chlorate; perchlorates, such as sodium perchlorate, potassium perchlorate, and ammonium perchlorate; inorganic peroxides, such as lithium peroxide, sodium peroxide, potassium peroxide, rubidium peroxide, cesium peroxide, magnesium peroxide, calcium peroxide, strontium peroxide, and barium peroxide; chlorites, such as sodium chlorite, potassium chlorite, copper chlorite, and lead chlorite; bromates, such as sodium bromate, potassium bromate, magnesium bromate, and barium bromate; nitrates, such as sodium nitrate, potassium nitrate, ammonium nitrate, barium nitrate, and silver nitrate; lodates, such as sodium iodate, potassium lodate, calcium lodate, and zinc lodate; permanganates, such as potassium permanganate, sodium permanganate, and ammonium permanganate; bichromates, such as sodium bichromate, potassium bichromate, and ammonium bichromate; periodates, such as sodium periodate; periodic acids, such as metaperiodic acid; chromium oxides, such as chromic anhydride (chromium trioxide); lead oxides, such as lead dioxide; fodine oxides, such as difodine pentoxide: nitrites, such as sodium nitrite and potassium nitrite; hypochlorites, such as calcium hypochlorite; chloroisocyanuric acids, such as trichloroisocyanuric acid; peroxodisulfates, such as ammonium peroxodisulfate; peroxoborates, such as ammonium peroxoborate; perchloric acid; hydrogen peroxide; nitric acid; halides, such as chlorine fluoride, bromine trifluoride, bromine pentafluoride, and iodine pentafluoride; iodine; and oxygen. These may be used alone or in combination as long as no danger is involved. It is also possible to use a compound which generates hydrogen peroxide in reaction with water or moisture in air, such as calcium dioxide. Among them, because of ease of reaction and high efficiency, preferred are sodium chlorate, sodium perchlorate, sodium peroxide, sodium chlorite, lead dioxide, hydrogen peroxide, calcium dioxide, and oxygen.

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[0072] When method (ii), in which a compound having at least two Isocyanato groups in each molecule is allowed to read with block copolymers, and thereby coupling of the block copolymers is performed via thiourethane bonds (NHCOS-), is employed, examples of the compound having at least two isocyanate groups in each molecule which may be used include, but are not limited to, diisocyanate compounds, such as hexamethylene diisocyanate, 2,4-to)lylene diisocyanate, prophene diisocyanate, 2,4-to)lylene diisocyanate, prophene diisocyanate, prophene diisocyanate, and tetramethylene diisocyanate, ethylene diisocyanate, methylenebic socyanate, prophene diisocyanate, methylenebic receptante, prophene diisocyanate, prophene diisocyanate, methylenebic prophene diisocyanate, diphenylmethane triisocyanate-modifications of these compounds formed by reaction of these compounds with polyhydric alcoholis; isocyanutare-modifications of these compounds; and polyvalent isocyanate compounds formed by reaction of these compounds with polyhydric alcoholis; isocyanutare-modifications of these compounds. These may be used alone or in combination. Almong them, in view of availability and reactivity, preferred are hexamethylene diisocyanate, 2,4-to)lylene diisocyanate, diphenylmethane diisocyanate, isophorone diisocyanate, xylylene diisocyanate, and bigocyanatemethylylcylohaxane.

[0073] In the reaction described above, a catalyst (urethane formation calalyst) may be used as necessary. For example, the catalysts cited in Polyurethanes: Chemistry and Technology, Part I, Table 30, Chapter 4. Saunders and Frisch, Interscience Publishers, New York, 1963 may be used in the sead-but usable catalysts are not limited thereto. As the urethane formation reaction catalysts which may be used in the reaction described above, the following catalysts are preferred because of their high activity: tin catalysts, such is tin catylate, this in stearate, dibutylitin discloter, dibutylitin bis(choxyairicate), dibutylitin oxide, dibutylitin bis(choxyairicate), dibutylitin oxide, dibutylitin bis(choxyairicate), di

10074] The amount of the catalyst added in the reaction is not particularly limited, but is preferably 0.0001 to 3 parts by weight, more preferably 0.001 to 0.5 parts by weight, and most preferably 0.003 to 0.1 parts by weight, based on 100 parts by weight of the block copolymer having mercaplo groups or mercaptide groups. If the amount is less than 0.0001 parts by weight, sufficient reactivity may not be obtained. If the amount exceeds 3 parts by weight, the properties of the resultant block copolymer, such as heter resistance, weatherability, and hydrolysis resistance, may be degraded. [0875] In method (iii) in which a compound having at least two isothiocyanato groups in each molecule is allowed to react with block copolymers, and thereby the block copolymers are coupled via dithiourethane bonds (AHCSS-), the compound having at teast two isothiocyanato groups in each molecule used is not particularly limited Examples thereof

include compounds obtained by replacing the isocyanato groups of the compound having at least two isocyanato groups in each molecule described above with isothicoyanato groups.

[9076] In method (iv) in which a compound having at least two unsaturated bonds in each molecule is added to block copolymers, and thereby the block copolymers coupled, examples of compounds having at least two unsaturated bonds in each molecule which may be used include, but are not limited to, butadiene, isoprene, chloroprene, 1,4-heptadiene, 1.5-hexadiene, 1.6-heptadiene, 1.7-octadiene, divinyl ether, diallyl ether, vinyl acrylate, vinyl methacrylate, allyl acrylate, allyl methacrylate, 1,2-diviny/benzene, 1,4-diviny/benzene, 1,3,5-triviny/benzene, bisphenol A divinyl ether, bisphenol A dialivi ether, trimethyloloropane divinyl ether, trimethyloloropane trivinyl ether, trimethyloloropane dialivi ether, trimethylolpropane Iriallyl ether, acrylate of alkyl-modified dipentaerythritol, ethylene oxide-modified bisphenol A diacrylate, acrylate of e-caprolactone-modified dipentaerythritol, diacrylate of caprolactone-modified neopentyl glycol hydroxypivalate, ditrimethylolpropane tetraacrylate, dipentaerythritol hexaacrylate, dipentaerythritol pentaacrylate, tetraethylene glycol diacrylate, tripropylene glycol diacrylate, trimethylolpropane ethylene oxide-modified triacrylate, trimethyloipropane triacrylate, neopentyl glycol diacrylate, diacrylate of neopentyl glycol hydroxypivalate, 1.9-nonanediol diacrylate, 1.4-butanediol diacrylate, 2-propenoic acid [2-f1.1-dimethyl-2-f(1-oxo-2-propenyl)oxylethyl]-5-ethyl-1.3-dioxane-5-vilmethyl ester, 1.6-hexanediol diacrylate, pentaerythritol triacrylate, ethylene glycol dimethacrylate, tetraethylene glycol dimethacrylate, triethylene glycol dimethacrylate, 1,3-butylene glycol dimethacrylate, trimethylolpropage trimethacrylate, pentagrythritol divinyl ether, pentagrythritol trivinyl ether, pentagrythritol diallyl ether, pentagrythritol ether, pentagr vthritoj triallyl ether, 1,4-butanediol divinyl ether, 1,4-butanediol diallyl ether, 1,5-pentanediol divinyl ether, 1,5-pentanediol dialityl ether, 1,6-hexanediol divinyl ether, 1,6-hexanediol dialityl ether, diethylene glycol divinyl ether, diethylene glycol dialiyl ether, tripropylene glycol divinyl ether, tripropylene glycol dialiyl ether, tetraethylene glycol divinyl ether, tetraethylene glycol dialfyl ether, poty(ethylene oxide) divinyl ether, poly(ethylene oxide) dialfyl ether, poly(propylene oxide) divinyl ether, poly(propylene oxide) diallyl ether, neopentyl glycol divinyl ether, neopentyl glycol diallyl ether, divinyl trimeilitate, trivinyl trimellitate, diallyl trimellitate, triallyl trimellitate, divinyl succinate, diallyl succinate, divinyl phthalate, diatity phthalate, divinyl maleate, diailyl maleate, divinyl terephthalate, diallyl terephthalate, divinyl carbonate. diallyl carbonate, 1,3,5-triacryloylhexahydro-1,3,5-triazine, triallyl isocyanurate, triallyl cyanurate, trimethallyl isocyanurate, furan, cyclopentadiene, dicyclopentadiene, and a maleimide compound represented by general formula (5):

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$$\begin{array}{c}
0 & 0 \\
N-R^5-N
\end{array}$$
(5)

(wherein R⁵ is a divalent organic group of 1 or more carbon atoms). In view of availability, preferably, R⁵ has 1 to 20 carbon atoms. These compounds may be used alone or in combination.

[0077] In method (v) in which dehydrocondensation is performed between a polyvalent carboxylic acid and block oppolymers, and thereby the block copolymers are coupled via thioester bonds, examples of polyvaient carboxylic acids which may be used include, but are not limited to, adipic acid, ilaconic acid, iminodiacetic acid, ethylenediaminetetraacetic acid, glutaric acid, citraconic acid, oxeracid acid, sabacid caid, 3.3° thiodipropionic acid, thiomalete acid, dodecanedioic acid, 1,2-cytolbexanediaminetetraacetic acid, brassylic acid, malonic acid, phitalic acid, isophithalic acid, aphthalenedicarboxylic acid, brassylic acid, malonic acid, phitalic acid, isophithalic acid, terpenthalic acid, aphthalenedicarboxylic acid, 5.5° thorough 2.4° they acid acid, aci

[9078] In method (vr) in which transesterification is performed between a polyvalent carboxylate ester and block copolymers, and thereby the block copolymers are coupled, as the polyvalent carboxylate ester, esters of the polyvalent carboxylate esters include, but are not limited to, methyl esters, ethyl esters, p-popyl esters, isopropyl esters, n-butyl esters, isobutyl esters, see-butyl esters, tert-butyl esters, decyl esters, isodecyl esters,

[0079] In method (vii) in which esterification is performed between a polyvalent carboxylic anhydride and block copolymers, and thereby the block copolymers are coupled, as the polyvalent carboxylic anhydrides of the polyvalent carboxylic acids described above may be used, but the anhydride which may be used is not limited thereto. These polyvalent carboxylic anhydrides may be used alone or in combination. In the reaction, any transesterification catalyst commonly used may be used. In this method, by removing water produced during reaction, the reaction can also be carried out effectively. As in the case described above, a method in which the resultant water is removed with a dehydrator, such as molecular sieves, a method in which the resultant water is removed by reaction with an orthocarboxylate ester or the like, or a method in which the resultant water is removed with an azeotropic solvent, such as toluren is a sopropriately employed.

[0080] In method (viii) in which dehydrohalogenation (acytation) is performed between a polyvalent carboxylic add halides and block copolymers, and thereby the block copolymers acoupled, as the polyvalent acyt halides include, but are not limited to, chlorinated compounds, such as succinyl dichloride, examples of polyvalent acyt halides include, but are not limited to, chlorinated compounds, such as succinyl dichloride, adypinyl dichloride, itaconyl dichloride, coxialy dichloride, transpired included, and the properties of the properties of the properties of the chlorine atoms in the above-mentioned chlorinated compounds by bromine atoms; and compounds formed by replacing the chlorine atoms in the above-mentioned chlorinated compounds by didne atoms. Among them, in view of availability and reactivity, preferred are chlorinated compounds. such as succinyl dichloride, maionyl dichloride, and fumaryl dichloride. These may be used alone or in combinetion. After the reaction, preferably, acids present in the system are removed by neutralization or distiliation under reduced pressure. If the acids are not removed, corrosion may occur.

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30 [3081] In method (ix) in which block copolymers are coupled by transesterification between a carbonate compound and the block copolymers, examples of carbonate compounds which may be used include, but are not limited to, dimethyl carbonate, diethyl carbonate, di-n-butyl carbonate, di-n-butyl carbonate, distorbutyl carbonate, distorbutyl

[0082] In method (x) in which a ketone is allowed to react with block copolymers to form thicketal bonds, and thereby the block copolymers are coupled, examples of ketones which may be used include, but are not limited to, acetylacetone, acetone, isophorone, disobutyl ketone, disopropyl ketone, cyclohexanone, cyclopentanone, 1,3-dihydroxyacetone dimethyl ether, 4,4-dimethoxy-2-butanone, diacetone acrylamide, diacetone alcohol, 4-hydroxy-2-butanone, methyl isobolyk ketone, methyl estyl es

[0063] In method (xi) in which block copolymers are coupled by dehydrocondensation between a compound having at least two hydroxyl groups in each molecule and the block copolymers, examples of compounds having at least two hydroxyl groups in each molecule which may be used include, but are not limited to, 3.6-dimethyl-4-cctyne-3.8-diol, 2.4-d'7-betramethyl-5-decyne-4.7-diol, 2.5-dimethyl-2-5-hexanethyl-5-hexanethyl-5-hexanethyl-5-hexanethyl-5-hexanethyl-1.5-hexanethy

thiodiglycol, trimethylolethane, trimethylolpropane, trimethylolpropane monoallyl ether, neopentyl glycol, 1.3-butanediol, 1.4-butanediol, 2-butyl-2-ethyl-1.3-propanediol, propylene glycol, dipropylene glycol, tripropylene glycol, 1.6-bexanediol, 1.2-6-bexanetriol, hexyfene glycol, pentaerythriol, 1.5-pentanediol, polyethylene glycol, polytropylene glycol, pentaerythriol, 1.5-pentanediol, polyethylene glycol, polytropylene glycol, 3-methyl-1, 5-pentanediol, catechol, 1.4-dihydroxyanthraquinone, 1.4-dihydroxyanthraquinone, 1.4-dihydroxyanthraquinone, 1.4-dihydroxyanthraquinone, 1.4-bitylorpylenethylen

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[9984] Among the coupling methods for coupling block copolymers having mercaptide groups, in method (xii) in which a compound having at least two halogen atoms in each molecule is allowed to react with block copolymers to form sulfide bonds (Williamson reaction), and thereby the block copolymers are coupled, examples of compounds having at least two halogen atoms in each molecule which may be used include, but are not limited to, methylene chloride, 1.1.1-trichloroethane, 1.2-dichloroethane, chloroform, trichloroethylene, tetrachloroethylene, 2,5-dichloroaniline, 3,4-dichloroaniline, 3,5-dichloroaniline, 2,4-dichlorobenzoic acid, 2,3-dichlorotoluene, 2,4-dichlorotoluene, 2,6-dichlorotoluene, 3,4-dichlorotoluene, 2,6-dichloro-4-nitroaniline, 1,4-dichloro-2-nitrobenzene, 2,4-dichloro-1-nitrobenzene, o-chlorobenzyl chloride, p-chlorobenzyl chloride, 2,6-dichlorobenzyl chloride, 3,4-dichlorobenzyl chloride. 2,3-dichlorobenzaldehyde, 2,4-dichlorobenzaldehyde, 2,6-dichlorobenzaldehyde, o-dichlorobenzene, m-dichlorobenzene. p-dichlorobenzene. 1.3.5-trichlorobenzene, 2.3-dichlorobenzoyl chloride, 2,4-dichlorobenzoyl chloride, 2.6-dichlorobenzovi chloride, carbon tetrachloride, 3,3'-dichloro-4,4'-diaminodiphenyimethane, 3,3'-dichloro-4,4'-diaminobiphenyl, 2.3-dichloro-5,6-dicyano-1,4-benzoquinone, 2,3-dichloro-1,4-naphthoquinone, 2,6-dichlorobenzal chloride, 2,8-dichlorobenzonitrile, octabromodiphenyl ether, 1,1,2,2-tetrabromoethane, 1,2-dibromo-4-(1,2-dibromoethyl)cyclohexane, 1,4-dibromobutane, 1,3-dibromopropane, 2,3-dibromo-1-propanol, 1,5-dibromopentane, decabromodiphenyl ether, tetradecabromo-p-diphenoxybenzene, tetrabromocyclooctane, tetramethylene chlorobromide, 2-(2-hydroxyethoxy)ethyl-2-hydroxypropyltetrabromophthalate, 1-bromo-2-chloroethane, 1-bromo-3-chloropropane, 1-bromo-6-chiorohexane, bromochloromethane, hexabromobenzene, pentamethylene chlorobromide, methylene didichloropentafluoropropane, 2,4-difluoroaniline, 2.6-difluorobenzonitrile, 2,2,3,3-tetrafluoropropanol, 2,2,2-trifluoroacetamide, trifluoroacetaldehyde hydrate, trifluoroethanol, trifluoroacetic acid, trifluoroacetic anhydride, trifluoroacetic acid ethyl ester, trifluoromethanesulfonic acid, trifluoromethanesulfonic acid anhydride, lithium trifluoromethanesulfonate, 2-(trifluoromethyl)benzaldehyde, 4-(trifluoromethyl)benzaldehyde, 2-(trifluoromethylibenzovi chloride, perfluorooctyl iodide, 2-perfluoroalkylethanol, perfluoroalkylethyl acrylate, perfluoropropyl vinyl ether, perfluoropolyalkenyl vinyl ether, 1,3-bis(trifluoromethyl)benzene, 1,4-bis(trifluoromethyl)benzene, 2,2-bis(4-hydroxyphenyl)hexafluoropropane, vinylidene fluoride, hexafluoroacetone trihydrate, hexafluoro-2-propanol, hexafluoropropylene, hexafluoropropylene oxide, 1,2-diiodoethane, and 1,4-diiodobenzene. These may be used alone or in combination. When transparency is required, produced salts are preferably removed by filtration or water washing. [0085] In method (xill) in which block copolymers are coupled by a reaction between a polyvalent carboxylic acid and the block copolymers, as the polyvalent carboxylic acid, the same polyvalent carboxylic acids as those used in method (v) may be used. The usable polyvalent carboxylic acids are not limited thereto. These may be used alone or in combination, in view of the durability of the resultant polymer, by-produced bases are preferably neutralized during

[0086] In method (xiv) in which block copolymers are coupled by a reaction between a polyvalent acyl halide and the block copolymers, as the polyvalent acyl halide, the same compounds as those used in method (xiv) in which disulfide bonds are formed between block copolymers, and thereby the block copolymers are coupled, as the oxidizing agent, the same compounds as those used in method (i) may be used.

[0088] When the block copolymers are coupled via the mercapto groups or mercaptide groups, it is also possible to use a compound heing the groups contained in the compounds described above. For example, compounds having carboxyl groups (method (v) and method (xiii)) and hydroxyl groups (method (xi)), such as selicylize acid and tactic acid: and compounds having halogen atoms (method (xii)) and carboxyl groups (method (v) and method (xiii), such as 4-chlorobearcic acid may be used to perform the coupling (reactions.

[0089] In order to carry out the coupling reactions efficiently, organic solvents may be used. Examples of organic solvents which may be used in the present invention include, but are not limited to, hydrocarbon solvents, such as heptane, octane, and mineral spirit: ester solvents, such as ethyl acetate, n-butyl acetate, isobutyl acetate, ethylene glycol monomouthyl ether acetate, extens solvents, such as acetone,

methyl ethyl ketone, methyl isobutyl ketone, dilsobutyl ketone, and cyclohexanone; ether solvents, such as teltrahydrofuran, diethyl ether, di-n-butyl ether, dioxane, ethylene glycol dimethyl ether, and ethylene glycol diethyl ether; and aromatic petroleum solvents, such as toluene, zylene, Swasoi 310 (manufactured by Cosmo Oil Co., Ltd.), Swasol 1000 (manufactured by Cosmo Oil Co., Ltd.), and Swasol 1500 (manufactured by Cosmo Oil Co., Ltd.). These solvents may be used alone or in combination. The reaction temperature is not particularly limited. In view of reactivity, the reaction temperature is preferably in the range of 0°C to 200°C.

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[9090] As the thermoplastic resin, which is contained in the thermoplastic resin composition of the present invention, various thermoplastic resins which are conventionally used may be used. Examples of thermoplastic resins include. but are not limited to, ionomer resins, such as SURLYN (manufactured by E. I. Du Pont de Nemours and Company), and HIMILAN (manufactured by Du Pont-Mitsui Polychemicals Co., Ltd.); polyacrylic acid hydrazide, isobutylenemaleic anhydride copolymers, acrytonitrile-styrene-acrytic rubber copolymers (AAS), acrytonitrile-EPDM-styrene copolymers (AES), acrylonitrile-styrene copolymers (AS), acrylonitrile-butadiene-styrene copolymers (ABS), and ABSvinyl chloride self-extinguishing resins, such as Kaneka Enplex (manufactured by Kaneka Corporation); ABS heatresistant resins, such as Kaneka MUH (manufactured by Kaneka Corporation); acrylonitrile-chlorinated polyethylenestyrene resins (ACS), methyl methacrylate-butadiene-styrene copolymers (MBS), ethylene-vinyl chloride copolymers, ethylene-vinyl acetate copolymers (EVA), modified ethylene-vinyl acetate copolymers, chlorinated ethylene-vinyl acetate copolymers, ethylene-vinyl acetate-vinyl chloride graft copolymers, ethylene-vinyl alcohol copolymers (EVOH), chlorinated poly(vinyl chloride), chlorinated polyethylene, chlorinated polypropylene, carboxyvinyl polymers, ketone resins, norbornene resins, polytetrafluoroethylene (PTFE), ethylene fluoride-propytene copolymers, tetrafluoroethylene-perfluoroalkyl vinyl ether copolymers (PFA), polychlorotrifluoroethylene, ethylene-tetrafluoroethylene copolymers, low-melting-paint ethylene-tetrafluoroethylene copolymers, poly(vinylidene fluoride) (PVDF), poly(vinylifluoride), polyacetal, polyamide 6, polyamide 66, polyamide 610, polyamide 612, polyamide 11, polyamide 12, copolymerized polyamides, polyamide MXD6, polyamide 46, methoxymethylated polyamides, polyamideimides, polyarylates, thermoplastic polyimides, polyether imides, polyether ether ketones, polyethylene, poly(ethylene oxide), poly(ethylene terephthalate) (PET), poly(ethylene naphthalate), poly(vinylidene chloride), poly(vinyl chloride) (PVC), polycarbonate, poly(vinyl acetate), polystyrene, polysulfone, poly(ether sulfone), poly(amine sulfone), polyparavinyiphenol, polyparamethylstyrene, polyallylamine, poly(vinyl alcohol) (PVA), polyvinyl ether, poly(vinyl butyral) (PVB), poly(vinyl formal) (PVF), polyphenylene ether, modified polyphenylene ether, poly(phenylene sulfide), polybutadiene, poly(butylene terephthalate) (PBT), polypropylene, polymethylpentene, poly(methyl methacrylate), and various types of liquid crystal polymers. These may be used alone or in combination. Among them, because of the excellence in heat resistance, weatherability, and oil resistance, at least one resin selected from the group consisting of poly(vinyl chloride), poly(methyl methacrviate), acrylonitrile-styrene copolymers, acrylonitrile-butadiene-styrene copolymers, polycarbonate, polyester resins, and polyamide resins is preferable.

[0091] An elastomer composition of the present invention contains the block copolymer and a synthetic rubber. Examples of such a synthetic rubber include, but are not limited to, styrene-butadiene rubber (SBR), butadiene rubber (BR), lesorene rubber (BR), elsorene rubber (BR), elsorene rubber (BR), esprene rubber (BR), esprene rubber (BR), esprene rubber (BR), esprene rubber (BR), exploitifile-butadiene copolymer rubber (BR), chloroprene rubber, butyl rubber (IR), urethane rubber, elsoren rubber, polysufider rubber, hydrogenated nitrile rubber, fluororubber, ethylene tetrafluoride-propylene-vinylidene fluoride rubber, acryfic rubber (ACM), chlorosufonated polyethylene rubber, epichlorohydrin rubber (OO), ethylene-acryfic rubber, norbormene rubber, styrene-based thermoplastic elastomers (TPU), polyseiz-based thermoplastic elastomers (TPU), polyseiz-based thermoplastic elastomers (TPU), polyseiz-based thermoplastic elastomers, (TPU), polyseiz-based thermoplastic elastomers, vinyl chloride-based thermoplastic elastomers (TPAE), 1,2-polybutadiene-based thermoplastic elastomers, vinyl chloride-based thermoplastic elastomers (TPAE), and fluorine-containing thermoplastic elastomers.

6 [0092] Each of the thermoplastic resin composition and the elastomer composition of the present invention may contain both a thermoplastic resin and a synthetic rubber besides the block copolymer. The composition may be dynamically crosslinked by a method which is commonly used.

[0093] In the thermoplastic resin composition and elastomer composition of the present invention, in order to adjust various physical properties, at least one type of additives may be compounded as necessary, besides the thermoplastic resin, the elastomer resin, and the block copolymer. As the additive, at least one material selected from the group consisting of plasticizers, thiotropy-improving agents, heat resistance-improving agents, stabilizers, antioxidents, ultraviolet absorbers, hindered amine light stabilizers (HALSs), antistatic agents, fire retardants, colorants, blowing agents, lubricants, mildewproofing agents, nucleating additives, vulcanization accelerators, aging resisters, vulcanizing agents, antiscorching agents, peptizers, tackifiers, latex coagulants, processing aids, inorganic fillers, and natural ruber may be used. Optimum additives may be selected depending on the types and compositions of the thermoplastic resins and the elastomers, the compositions of the block copolymers, the applications of the compositions, etc.

(1004) The black polymers of the present invention have heat resistance, weatherability, oil resistance, and flame retardancy which are originally exhibited by the acrylonitrile and methacrylonitrile polymers, and furthermore are ex-

cellent in low-temperature resistance. Consequently, the block copolymers can be used in various applications, such as films, sheets, tapes, hoses, tubes, gaskets, packings, grips, various molded objects, sealants, damping materials, pressure-sensitive adhesives, adhesives, resin modifiers, coaling materials, potting materials, textiles, materials for thermoplastic resin compositions, and materials for elastomers.

[0095] The thermoplastic resin compositions of the present invention contain the block copolymers and thermoplastic resins. The elastomer compositions of the present invention contain the block copolymers and synthetic rubbers. These compositions can be used widely in various applications, such as films, sheets, tapes, hoses, tubes, gaskets, packings, grips, containers, various molided objects, sealants, damping materials, pressure-sensitive adhesives, adhesives, coating materials, potting materials, and textiles.

Best Mode for Carrying Out the Invention

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[9996] While the present invention will be described based on the examples below, it is to be understood that the invention is not limited thereto.

[0097] In the examples below, the weight-average molecular weight (Mw), number-average molecular weight (Mh), and molecular weight distribution (MwMh) were determined by gel permeation chromatography (GPC). In the GPC, chloroform, letrahydrofuran, or dimethytformamide was used as an eluent, and a polystyrene gel column was used. The analysis was carried out on the basis of a polystyrene standard sample.

20 (Example 1; Synthesis of mercapto group-terminated acrylonitrile-n-butyl acrylate diblock copolymer)

[0098] Into a 2 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was piaced 490 g of distilled water and 0.56 g of sodium dodecyl sulfate, and the reactor was nitrogen-purged while the reaction mixture was being stirred at 80°C. A mixed solution of 8.8 g of acrylonittile and 1.09 of a compound represented by formula (6):

$$\begin{array}{c}
S & CH_3 \\
C-S-C & CH_3
\end{array}$$
(6)

was added into the reactor, and stirring was performed at 80°C for 20 minutes. Next, 0.93 g of 4.4"-azxbis(4-cyanovaleric acid) logether with 2.5 g of distilled water was added into the reactor. Stirring was performed at 80°C for 30 minutes, and then 45.0 g of acrylontifie was dripped from the dropping lunnel for over 1 hour. After dripping was performed at 80°C for 5 hours, and sampling was performed. Production of polyacrylonitrile (Mw = 13.700, Mn = 10.300, and Mw/Mn = 1.33) was confirmed by gel permeation chromatography analysis.

[0099] Next, 20.0 g of n-butyl acrylate was added into the reactor, and 0.40 g of 4.4"-azobis(4-cyanovaleric acid) together with 10 g of distilled water was further added thereinto. Stirring was performed at 80°C for 1 hour, and then 80.0 g of n-butyl acrylate was dipped from the dropping furnel for over 2 hours. After dripping was completed, the mixture was stirred at 80°C for 5 hours and then cooled to room temperature. A salting-out method was performed, followed by filtration, washing, and drying. Thereby, production of an acrylonitrile-n-butyl acrylate diblock copolymer (Mw = 48,500, Mn = 34,500, and Mw/Mn = 1.41) was confirmed.

[0100] The diblock copolymer (50 g) was dissolved in 200 mL of toluene, and 15 g of monoethylamine was added thereinto, followed by stifring at 30°C for 8 hours. '14 Mift? analysis and IR analysis confirmed that the thiocarbonyithio groups at ends were quantitatively converted into mercapto groups. The toluene solution was washed with water and then pound into methanol to precipitate and isolate the block copolymer.

(Example 2: Synthesis of mercapto group-terminated methacrylonitrile-n-butyl acrylate diblock copolymer)

[0101] Into a 2 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was placed 490 g of distilled water and 0.55 g of sodium dodecyl sulfate, and the reactor was nitrogen-purged while the reaction muture was being stirred at 80°C. A mixed solution of 11.3 g of methacrylonitrile and 1.88 g of a compound represented by formula (6):

$$\begin{array}{c|c}
S & CH_3 \\
C-S-C & CH_3
\end{array}$$
(6)

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was added into the reactor, and stirring was performed at 80°C for 20 minutes. Next, 0.83 g of 4,4-azobis(4-cyanovaleric acid) together with 25 g of distilled water was added into the reactor. Stirring was performed at 80°C for 30 minutes, and then 57.7 g of methacrylonitrile was dripped from the dropping funnel for over 1 hour. After dripping was completed, stirring was performed at 80°C for 5 hours, and sampling was performed. Production of polymethacrylonitrile (Mw = 18 400. Mn = 13.000. and Mw/Mn = 1.26) was confirmed by one permeation chromatography analysis.

[0192] Next, 20.0 g of n-butyl acryster was added into the reactor, and 0.36 g of 4.4-axobis4(4-yanovaleric acid) together with 1.3 g of distilled water was further added thereinto. Stirring was performed at 80°C for 1 hour, and then 80.0 g of n-butyl acryster was dripped from the dropping funnel for over 2 hours. After dripping was completed, the mixture was stirred at 80°C for 4 hours and then cooled to norm temperature. A satting-out method was performed, followed by filtration, washing, and drying. Thereby, production of a methacrylonitrilen-butyl acrysted blobck copolymer

(Nw = 49,900, Mn = 36,300, and Mw/Mn = 1.37) was confirmed.

[0103] The diblock copolymer (70 g) thus obtained was dissolved in 300 mL of folluene, and 18 g of monoethylamine was added thereinto, followed by stirring at 30°C for 16 hours. ¹H NMR analysis and IR confirmed that the thiocarbonythio groups at ends were quantitatively converted into mercapto groups. The tolluene solution was washed with water and then pourcel into methanol to precipitate and isolate the block copolymer.

(Example 3: Synthesis of mercapto group-terminated acrylonitrile-(n-butyl acrylate/ethyl acrylate) diblock copolymer)

[0104] Into a 300 mL reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was placed 110 mg of sodium dodecyl sulfonate and 100 g of distilled water, and the reactor was nitrogen-purged while the reaction mixture was being stirred at 80°C. A compound (217 mg) represented by formula (6):

$$\begin{array}{c}
S & CH_3 \\
C-S-C & CH_2
\end{array}$$
(6)

which was dissolved in 1.6 g of acrylionitrile, was added into the reactor, and after 20 minutes, 185 mg of 4.4"-azobis (4-cyanovaleric acid) together with 5 g of distilled water was added into the reactor. Suring was performed a 16°C for 20 minutes, and then 9.3 g of acrylionitrile was dripped from the dropping tunnel for over 50 minutes. After dripping was completed, stirring was performed at 80°C for 3 hours, and sampling was performed. Production of polyacrylionitrile (Mw = 18,100, M = 12,700, and MeW/M = 1.42) was confirmed by get permeation chromatography analysis.

[0105] Next, a mixed solution of 10.0 g of n-butyl acrylate and 7.8 g of ethyl acrylate was dripped from the dropping funnel for over 1 hour. Stirring was performed at 80°C for 5 hours, and an emulsion was thereby prepared. Sampling was performed and production of a thiocarbonylthio group-terminated acrylonitrile-(n-butyl acrylate/ethyl acrylate) diblock copolymer (Mw = 48.500, Mn = 31,600, and Mw/Mn = 1.53) was confirmed.

[0106] Diethylamine (20 g) was added into the emulsion, and stirring was performed at 60°C for 5 hours. A sattingout method was performed, followed by filtration, washing, and drying, and a polymer was thereby produced. It was confirmed that the polymer was a mercapto group-terminated acrylonitrile-(n-butyl acrylate/ethyl acrylate) diblock copolymer (Mw = 48,700, Mn = 31,300, and Mw/Mn = 1.56). (Example 4: Synthesis of mercaptide group-terminated (acrylonitrile/methyl methacrylate)-{n-butyl acrylate/ 2-hydroxyethyl acrylate) diblock copolymer)

[0107] Into a 1 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was placed 200 mg of sodium dodecyl sulfonate and 200 g of distilled water, and reactor was nitrogen-purged while the reaction mixture was being stirred at 80°C. A compound (651 mg) represented by formula (6).

$$\begin{array}{c}
S & CH_3 \\
C-S-C & CH_3
\end{array}$$
(6)

which was dissolved in 5.0 g of methyl methacrylato, was added into the reactor, and after 20 minutes, 500 mg of 4.4" acabis(4-cyanovaleric acid) together with 12 g of distilled water was added into the reactors. Stirring was performed at 80°C for 30 minutes, and then a mixed solution of 18.6 g of acrylonitrile and 6.8 g of methyl methacrylate was cripped from the dropping funnel for over 1 hour. Stirring was then performed at 80°C for 6 hours, and sampling was performed. Production of a thiocarbonylithio group-terminated acrylonitrile/methyl methacrylate random copolymer was confirmed. [9108] Next, 200 mg of 4.4"-szobis(4-cyanovalence acid) was added into the reactor, and a mixed solution of 50.0 g of hourly acrylate was 18.9 g of 24-hydroxyethyl acrylate of 18.0 g of 18.0 minutes. Stirring was performed at 80°C for 5 hours, and the resultant emulsion was sampled. Production of a thiocarbonylithio group-terminated (acrylonitrile/methyl methacrylate)-(houtyl acrylate/2-hydroxyethyl acrylate) billoick copolymer was confirmed. The emulsion was saited out, followed by filtration, washing, and drying, and the diblock copolymer was thereby prepared.

[0108] The diblock copolymer (30 g) thus obtained was dissolved in 100 mL of dehydrated foluene, and 4 g of sodium methylate was added thereinto, followed by stirring at 50°C for 2 hours. A foluene solution containing a mercapitide group-terminated (acrylonitrie/methyl methacrylate)-(n-butyl ecrylate/2-hydroxyethyl acrylate) diblock copolymer was thereby prepared. A portion of the solution was collected as a sample and hydrolyzed, and then poured into hoxane to precipitate and isolate the copolymer. 'H NMR analysis and GPC analysis confirmed the production of a mercapito group-terminated (acrylonitrile/methyl methacrylate)-(n-butyl acrylate/2-hydroxyethyl acrylate) diblock copolymer (Mw = 52,100. Mn = 35,200, and Mw/Mn = 1.482.)

(Example 5; Synthesis of acrylonitrile-n-butyl acrylate-acrylonitrile triblock copolymer)

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(0110) Toluene (150 parts by weight) and lead dioxide (0.5 parts by weight) were added to the mercapto group-terminated acrylonitrile-n-butyl acrylate diblock copolymer (100 parts by weight) synthesized in Example 1, and thorough mixing was performed. The mixture was placed into a slab mold and dried in an air atmosphere at 80°C for 15 hours. 1H NMR analysis, iR analysis, and gel permeation chromatography analysis confirmed that the resultant sheet-shaped polymer was an acrylonitrile-n-butyl acrylate-acrylonitrile triblock copolymer having disulfide bonds in the main chain (Mw = 101,600, Mn = 67.300, and Mw/M = 1.51).

(Example 6: Synthesis of acrylonitrile-n-butyl acrylate-acrylonitrile triblock copolymer)

[0111] Dehydrated toluene (200 parts by weight), hexamethylene diiscoyanate (0.24 parts by weight), and dibutytiin bis(isooch)t thiogiycolate) (0.001 parts by weight) were added to the mercapto group-terminated acrylonitrilen-butyl acrylate diblock copolymer (100 parts by weight) synthesized in Example 1, and string was performed at 80°C for 10 hours. The solvent was removed by distillation. ¹H NMR analysis, IR analysis, and gel permeation chromatography analysis confirmed that the resultant polymer was an acrylonitrilen-butyl acrylate-acrylonitrile triblock ecoplymer having thiourethane bonds in the main chain (Ahw = 99,700, Ma = 65,00, and MwMm = 1.50).

55 (Example 7: Synthesis of methacrylonitrile-n-butyl acrylate-methacrylonitrile triblock copolymer)

[0112] Calcium dioxide (1 part by weight) was added to the mercapto group-terminated methacrylonitrile-n-butyl acrylate diblock copolymer (100 parts by weight) synthesized in Example 2, and the mixture was roll-kneaded at 100°C

and formed into a sheet. Furthermore, the sheet was heated at 100°C for 5 hours and matured at room temperature for 3 days, 1H NMR analysis, IR analysis, and get permeation chromatography analysis confirmed that the resultant sheet-shaped polymer was a methacrylonitrile—butyl acrylate—methacrylonitrile triblock copolymer having disulfide bonds in the main chain (Mw = 102,700, Mn = 71,400, and Mw/Mn = 1.44).

(Example 8: Synthesis of methacrylonitrile-n-butyl acrylate-methacrylonitrile triblock copolymer)

[0113] Dehydrated toluene (100 parts by weight), isophorone diisocyanate (0.31 parts by weight), and dibutyniti bisacetylacetonate (0.001 parts by weight) were added to the mercapto group-terminated methacytonitrilen-butyl acrylate diblock copolymer (100 parts by weight) synthesized in Example 2, and the mixture was stirred at 80°C for 8 hours. Toluene was then removed by distillation. ¹H NMR analysis, IR analysis, and gel permeation chromatography analysis confirmed that the resultant polymer was a methacrylonitrilen-butyl acrylate-methacrylonitrile triblock copolymer having thiourethane bonds in the main chain (May = 85.00, Mn = 69.900, and Mw/MM = 1.41).

15 (Example 9: Synthesis of acrylonitrile-(n-butyl acrylate-ethyl acrylate)-acrylonitrile triblock copplymen)

[9144] Toluene (200 parts by weight) was added to the mercapto group-terminated acrylonitrile-(h-butyl acrylate) ethyl acrylate) diblock copolymer (100 parts by weight) synthesized in Example 3, and the mixture was stirred at 90 for 18 hours under blown air. Toluene was removed by distillation. "H NMR analysis, IR analysis, and get permeation chromatography analysis confirmed that the resultant polymer was an acrylonitrile-(n-butyl acrylate)acrylonitrile triblock copolymer having disulfide bonds in the main chain (Mw = 102.400, Mn = 61,100, and Mw/Mn = 1.68).

(Example 10: Synthesis of acrylonitrile-(n-butyl acrylate/ethyl acrylate)-acrylonitrile triblock copolymer)

[0115] Toluene (200 parts by weight) and diphenylmethane dilsocyanate (0.4 parts by weight) were added to the mercapto group-terminated acrylonitrile-(n-butyl acrylate) acrylate) diblock copolymer (100 parts by weight) synthesized in Example 3, and the mixture was stirred at 100°C for 8 hours. Toluene was removed by distiliation. 1H MMR analysis, IR analysis, and gel permeation chromatography analysis confirmed that the resultant polymer was an acrylonitrile-(n-butyl acrylate/ethyl acrylate/bap-acrylonitrile triblock copolymer having thiourethane bonds in the main chain (Mw = 10.70, Mn = 60,900, and Mw/Mn = 1.65).

(Example 11: Synthesis of (acrylonitrile/methyl methacrylate)-(n-butyl acrylate/2-hydroxyethyl acrylate)-(acrylonitrile/methyl methacrylate) triblock copolymer)

[0116] To a toluene solution containing the (acrylonitrile/methyl methacrylate)-(in-butyl acrylate)-thydroxyethyl acrylate) diblock copolymer (100 parts by weight), and the basis of the polymer) was added 1,2-dicthoroethane (0.14 parts by weight), and the mixture was stirred at 80°C for 20 hours. The resultant toluene solution was washed with water and dried, and the solvent was removed by distillation. HI NMR analysis and GPC analysis confirmed that the resultant polymer was an (acrylonitrile/methyl methacrylate)-(hebutyl acrylate2-lydroxyethyl acrylate)-(acrylonitrile/methyl methacrylate) triblock copolymer having sulfide bonds (Nw = 101,700, Mm = 65,100, and MwMMn = 1.65).

(Example 12: Synthesis of mercapto group-terminated acrylonitrile-n-butyl acrylate diblock copolymer)

45 [0117] Into a 2 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was piaced 490 g of distilled water, 20 g of polyoxyethylene nonylphenyl ether as a nonionic american sitier, and 2 g of hexadecane as an emulsifying aid, and the reactor was nitrogen-purged while the reaction mixture was being stirred at 80°C. A mixed solution of 8.9 g of acrylonitrite and 1.1 g of a compound represented by formula (6):

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$$\begin{array}{c}
S & CH_3 \\
C-S-C & CH_3
\end{array}$$
(6)

was added into the reactor, and stirring was performed at 70°C for 30 minutes. Next, 0.92 g of 4,4'-azobis(4-cyanovaleric acid) together with 25 g of distilled water was added into the reactor. Stirring was performed at 70°C for 1 hour, and then 45 g of acryfonitfile was dripped from the dropping funnel for over 1 hour. As soon as dripping was completed, the temperature was raised to 80°C and stirring was performed for 5 hours. Sampling was then performed. Production of polyacryfonitrile (Mw = 13.000, Mn = 10,000, and Mw/Mn = 1.30) was confirmed by gel permeation chromatography analysis.

[0118] Next, 20 g of n-butyl acrylate was added into the reactor, and 0.25 g of 4,4'-azobis(4-cyanovaleric acid) together with 5 g of distilled water was further added thereinto. Stirring was performed at 80°C for 1 hour, and then 80 g of n-butyl acrylate was dripped from the dropping funnel for over 2 hours. After dipping was completed, the mixture was stirred at 80°C for 6 hours and then cooled to room temperature. A salting-out method was performed, followed by filtration, washing, and drying. Thereby, production of an acrylonitrile-n-butyl acrylate diblock copolymer (Mw = 47.500, Mn = 34,000, and thw/Mn = 1.40) was confirmed.

[0119] The diblock copolymer (50 g) thus produced was dissolved in 200 mL of toluene, and 20 g of diethylamine was added thereinto, followed by stirring at 50°C for 8 hours. The reaction solution was purified with a silica gel column, and toluene was removed by distillation. ¹H NMR analysis and IR analysis confirmed that the thiocarbonylthic groups at ends were quantitatively converted into mercapto groups.

(Example 13: Synthesis of acrylonitrile-n-butyl acrylate-acrylonitrile triblock copolymer)

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[0120] Toluene (100 parts by weight), dimethylformamide (50 parts by weight), and lead dioxide (0.5 parts by weight) were added to the mercapto group-terminated acrylonitrile-n-butyl acrylate diblock copolymer (100 parts by weight) synthesized in Example 12, and thorough mixing was performed. The mixture was placed into a slab moid and dried in an air atmosphere at 80°C for 5 hours. Deseration was performed under reduced pressure at 80°C to remove the solvent, and healting was further performed in an air atmosphere at 80°C for 10 hours. ¹H NIMR analysis, if analysis, and get permeation chromatography analysis confirmed that the resultant polymer constituting the sheet was an acrylonitrile-n-butyl acrylate-acrylonitrile triblock copolymer having disulfide bonds in the main chain (Mw = 99,900, Mn = 66,500, and Mw/Mn = 15.0).

40 (Production Example 1: Synthesis of mercapto group-terminated polyacrylonitrile)

[0121] Into a 2 it, reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was piaced 490 g of distilled water and 0.56 g of sodium dodecyl sulfate, and the reactor was nitrogen-purged while the reaction mixture was being stirred at 80°C. A mixed solution of 8.8 g of acrylonitrile and 1.09 g of 2-(2-phenylpropyl) disthibotenzoate represented by formula (6):

$$\begin{array}{c}
S & CH_3 \\
-C-S-C-
\end{array}$$
(6)

was added into the reactor, and stirring was performed at 80°C for 15 minutes under nitrogen flow. Next. 0.93 g of 4.4°azobis(4-cyanovaleric acid) together with 25 g of distilled water was added into the reactor. Stirring was performed a 80°C for 1 hour, and then 45.0 g of acrylonitrile was dripped from the dropping funnel for over 2 hours. After dripping was completed, stirring was further performed at 80°C for 5 hours. The resultant emulsion was cooled to 30°C, and 30 g of monoethylamine was added thereinto, followed by stirring at 30°C for 10 hours. The emulsion was satted out, followed by filtration and washing. Polyacrylonitrile having a mercapto group at one end was thereby produced (Mw = 13,400, Mn = 10,800, and Mw/Mn = 1,24).

(Production Example 2: Synthesis of mercapto group-terminated acrylonitrile-methyl methacrylate random copolymer)

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[0122] Into a 1 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was placed 200 mg of sodium dodecyl sulfonate and 200 g of distilled water, and the reactor was nitrogen-purged while the reaction mixture was being stirred at 80°C. 2-(2-Phenylpropyl) dithiobenzoate (651 mg) represented by formula (6):

$$\begin{array}{c|c}
S & CH_3 \\
\hline
C-S-C & CH_2
\end{array}$$
(6)

was dissolved in a mixed solution of 18.6 g of acrylonitrile and 11.8 g of methyl methyarylate, and the solution was dripped from the dropping funnel at one time. After 20 minutes, 500 mg of 4.4-azobie(4-y-anovaleric acid) together with 1.2 g of distilled water was added into the reactor. Stirring was performed at 80°C for 4 hours, and the reaction mixture was cooled to 30°C. Monoethylamine (10 g) was added thereinto, and stirring was performed at 30°C for 10 hours. The resultant emulsion was salted out, followed by filteration and washing. An acrylonitrile-methyl methacrylate random copolymer having a mercapto group at one end was thereby produced (Mw = 11,100. Mn = 9,700, and Mw/Mn = 1.14).

(Production Example 3: Synthesis of poly(n-butyl acrylate) having mercapto group at one end)

[0123] Into a 2 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was placed 0.56 g of sodium dodecyl suifate and 490 g of distilled water, and the reactor was nitrogen-purged while the reaction mixture was being stirred at 80°C. 2-(2-Phenylpropyl) dithiobenzoate (1.09 g) represented by formula (6):

$$\begin{array}{c}
S & CH_3 \\
C-S-C & CH_2
\end{array}$$
(6)

was dissolved in 16.5 g of n-butyl acrylate, and the solution was dripped from the dropping funnel at one time. After 20 minutes, 0.93 g of 4.4"-ezobis(4-cyanovaloria exid) together with 25 g of distilled water was added into the reactor. Stirring was performed at 80°C for 30 minutes, and then 100 g of n-butyl acrylate was dripped from the dropping funnel for over 2 hours. Stirring was further performed at 80°C for 5 hours, and the reaction mixture was cooled to 30°C. Monoethylamine (30 g) was added thereinto, and stirring was performed at 30°C for 10 hours. The resultant emulsion was safted out, followed by filtration and washing. Poly(n-butyl acrylate) having a mercaplo group at one end was thereby produced (flw = 38.90.0, Mn = 30.900, and Mw/Mn = 1.28).

(Production Example 4: Synthesis of poly(n-butyl acrylate) having mercapto groups at both ends)

[0124] into a 1 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, and a reflux condenser

tube, was placed 181 g of n-butyl acrylate, 40 mg of 1,1'-azobis(1-cyclohexanecarbonitrile), 635 mg of 1,4-bis(thioben-zovlthiomethy/)benzene represented by formula (7):

and 300 mL of toluene, and the reactor was nitrogen-purged. The reaction liquid was heated at 90°C for 5 hours while being stirred. Sampling was performed, and GPC analysis confirmed the production of a polymer (Mw = 77,000, Mn = 56,900, and Mw/Mn = 1.35). "In NMR measurement confirmed that thiocarbony/thio groups were introduced into both ends of poly(n-butyl acrylate), and the introduction rate was 93% on the both-ends basis. The conversion rate of nbutyl acrylate was 55%.

[0125] Monoethylamine (30 g) was added into the resultant toluene solution containing poly(n-butyl acrylate) having thiocarbonyithio groups at both ends, followed by stirring at 30°C for 5 hours. Sampling was performed, and 1H NMR measurement confirmed the production of poly(n-butyl acrylate) having mercapto groups at both ends. The introduction rate of mercapto groups was 90% on the both-ends basis.

(Production Example 5: Synthesis of n-butyl acrylate-2-methoxyethyl acrylate random copolymer)

[0126] Into a 1 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a reflux condenser, and a dropping funnel, was placed 410 mg of sodium dodecyl sulfonate and 400 g of distilled water, and the reactor was nitrogen-purged while the reaction mixture was being sittered at 80°C. A compound (23.34 g) represented by formula (8):

which was dissolved in 50 g of n-butyl acrylate, was added into the reactor, and stirring was performed at 80°C for 20 minutes under introgen flow, Next. 7.0 g of 4.4-zea/bis4(~eya-proveleric acid) together with 25 g of distilled water was added thereinto. Stirring was performed at 80°C for 30 minutes, and then a mixed solution of 100 g of n-butyl acrylate and 50 g of 2-methoxyethyl acrylate was dripped from the dropping funnel for over 1.5 hours. After dripping was concleted, the mixture was stirred at 80°C for 4 hours, and the resultant emulsion was then cooled to room temperature. A satting-out method was performed, followed by filtration and washing. Thereby, an n-butyl acrylate-2-methoxyethyl acrylate random copolymer having thiocarbonythiol groupus at both ends was produced. GPC analysis and "H MMR analysis confirmed that in the polymer, Mw = 4,320. Mn = 3,970, and Mw/Mn = 1.09 and that the introduction rate of thiocarbonythiol groups was 97% on the both-ends basis.

[0127] The polymer having thiocarbonylthio group at both ends (180 g) was dissolved in 200 mL of toluene, and 20 g of monoethylamine was added thereinto, followed by stirring at 6°C for 10 hours. By removing excess monoethylamine and toluene, an n-butyl acrylate-2-methoxyethyl acrylate random copolymer having mercapto groups at both ends was produced.

50 (Production Example 6: Synthesis of mercapto group-terminated poly(viny) chloride))

[0128] Into a 300 L stainless steel autoclave was placed 130 kg of ion-exchanged water, 100 kg of viryl chloride monomer, 500 g of 2,2-azobis(isobutylvaleronitrie). 400 g of sodium lauryl sulfate, and 110 g of a compound represented by formula (9) below.

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$$\begin{array}{ccc}
& & & \\
N - C - S - CH_2CN & & (9)
\end{array}$$

[0129] Homogenizing was performed for 90 minutes with a homogenizer, and polymerization was then carried out at 50°C. When the internal pressure of the reactor became lower than the saturated vapor pressure of the viny chloride monomer at 50°C by 14 g/cm², urreacted monomers were removed from the reactor by distillation, A vinyl chloride polymer latex was thereby produced. Excess vinyl chloride was removed, and the reactor was filled with ammonia at a rate of 5 kg/cm² (20°C). Stirring was performed at 50°C for 5 hours. By spray drying the emulsion, mercapto group-terminated polyfvinier this forther's was produced (fixe = 61.000, Mm = 41.000, and Mw/mlh = 14.9).

(Production Example 7; Synthesis of mercapto group-terminated polyacrylonitrile)

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[0130] Into a 2 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was placed 490 g of distilled water, 20 g of polyethylene glycol nonylphenyl ether as a no indicated and a sane mulasifying aid, and the reactor was nitrogen-purged while the reaction mixture was being siltred at 80°C. A mixed solution of 8.9 g of acrylonitrile and 1.1 g of a compound represented by formula (6):

$$\begin{array}{c}
S & CH_3 \\
C-S-C & CH_3
\end{array}$$
(6)

was added into the reactor, and stirring was performed at 70°C for 30 minutes. Next, 0.92 g of 4.4-azobis(4-cyanovaleric acid) together with 25 g of distilled water was added into the reactor. Stirring was performed at 70°C for 1 hour, and then 45 g of acrylonititile was dripped from the dropping lunnel for over 1 hour. As soon as dripping was completed, the temperature was raised to 80°C and stirring was performed for 6 hours. The nikrure was cooled to room temperature, and an emulsion was greated. The emulsion was dried and washed with acetone, and then reprecipilation was performed with dimethylformamide-acetone. Polyacrylonitrile was thereby produced. Gel permeation chromatography analysis confirmed that Mv = (13,100, Mn = 9,800, and Mw/Mm/Mn = 1.34.

[0131] The resultant polyacrytonitrile (30 g) was dissolved in 300 mL of dimethylformamide, and 8 g of diethylamine was added thereinto, followed by stirring at 50°C for 16 hours. The reaction solution was concentrated and reprecipitated with methanol. Mercapto group-terminated polyacrytonitrile was thereby produced. NMR analysis confirmed that the introduction rate of mercapto groups was 91% on the single-end basis.

45 (Example 14: Synthesis of acrylonitrile-n-butyl acrylate diblock copolymer)

[0132] The poly(n-buly) acrylate) (286 parts by weight) synthesized in Production Example 3, toluene (300 parts by weight), and lead dioxide (0.5 parts by weight) were added to the mercapto group-terminated polyacrylonitrile (100 parts by weight) synthesized in Production Example 1, and the mixture was stirred at 80°C for 10 hours in an air atmosphere. ¹H NMR analysis confirmed that an acrylonitrile-n-bulyl acrylate diblock copolymer in which coupling was performed via disullide bonds was produced. GPC analysis confirmed that Mw = 81,400, Mn = 45,400, and Mw/Mn = 1,35.

(Example 15: Synthesis of acrylonitrile-n-butyl acrylate-acrylonitrile triblock copolymer)

[0133] In a nitrogen atmosphere, dehydrated toluene (300 parts by weight), isophorone discocyanate (0.70 parts by weight), and dibutyllin bis(isooctyl thiogiycolate) (0.05 parts by weight) were added to the poly(n-butyl acrylate) having mercapto groups at both ents (100 parts by weight) synthesized in Production Example 4, and the mixture was stirred

at 80°C for 8 hours to synthesize poly(n-butyl acrylate) having isocyanato groups at both ends. The mercapto groupterminated polyacrylonitrile (34 parts by weight) synthesized in Production Example 1 was added thereinto, and stirring was performed at 80°C for 16 hours. 14 hMR and GPC analysis confirmed that an acrylonitrilen-butyl acrylate-acrylonitrile triblock copolymer was produced (Mw = 127,000, Mn = 84,500, and Mw/Mn = 1.50).

(Example 16: Synthesis of acrylonitrile-(n-butyl acrylate/2-methoxyethyl acrylate)-acrylonitrile triblock copolymer)

[0134] In a nitrogen atmosphere, dehydrated toluene (300 parts by weight), isophorone diisocyanale (7.24 parts by weight), and dibutyltin bis(isooctyl thioglycolate) (0.05 parts by weight) were added to the n-butyl acrylate-2-methox-yethyl acrylate random copolymer having mercapto groups at both ends (100 parts by weight) synthesized in Production Example 5, and the mixture was stirred at 80°C for 10 hours to synthesize an n-butyl acrylate/2-methox-septily acrylate random copolymer having isocyanato groups at both ends. The mercapto group-terminated polyacrylonitrile (176 parts by weight) synthesized in Production Example 1 was added thereinto, and stirring was performed at 80°C for 16 hours. ¹H NMR and GPC analysis confirmed that an acrylonitrile-(n-butyl acrylate/2-methoxyethyl acrylate)-acrylonitrile triblock copologymer was produced MW = 76.700. Mn = 43.600, and MW/MM = 1.76).

(Example 17: Synthesis of acrylonitrile-vinyl chloride diblock copolymer)

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[0135] The mercapto group-terminated poly(vinyl chloride) (380 parts by weight) synthesized in Production Example 6 and lead dioxide (1 part by weight) were added to the mercapto group-terminated polysorytonitrile (100 parts by weight) synthesized in Production Example 1, and the mixture was meti-kneaded with a twin-screw extruder (finiet temperature 80°C; outlet temperature 160°C) and formed into a strand. The resultant strand was cut into pellets. The pellets were dried by heating at 80°C for 30 hours in an air atmosphere. 14 hMR and GPC analysis confirmed that an acrylonitrie-vinyl chloride diblock copolymer was produced (Mw = 71,100, Mn = 49,400, and Mw/Mn = 1,44).

(Example 18: Synthesis of (acrylonitrile/methyl methacrylate)-n-butyl acrylate diblock copolymer)

[0136] In a nitrogen atmosphere, dehydrated toluene (300 parts by weight), hexamethylene diisocyanate (5.78 parts by weight), and dibutylin big(isoocty) thioglycolate) (0.05 parts by weight) were added to the mercapto group-terminated acrylonitifial/methyl methacrylate random copolymer (100 parts by weight) synthesized in Production Example 2, and the mixture was stirred at 80°C for 10 hours to produce an isocyanato group-terminated acrylonitrifie/methyl methacrylate random copolymer. The mercapto group-terminated poly(n-butyl acrylate) (105 parts by weight) synthesized in Production Example 3 was added thereinto, and stirring was performed at 100°C for 8 hours in a nitrogen atmosphere. 1H NMR and GPC enarysis confirmed that an (acrylonitripliently) methacrylate)-n-butyl acrylate diblock copolymer was produced (Mw = 62,900, Mn = 45,200, and Mw/Mn = 1.39).

(Example 19: Synthesis of (acrylonitrile/methyl methacrylate)-n-butyl acrylate-(acrylonitrile/methyl methacrylate) triblock copolymer)

40 [0137] In a nitrogen atmosphere, dehydrated toluene (500 parts by weight), lsophorone diisocyanate (0.70 parts by weight), and diutyltin bis(socot) thioglycotate) (0.05 parts by weight) were added to the poly(n-butyl acrylate) having mercapto groups at both ends (100 parts by weight) synthesized in Production Example 4, and the mixture was stirred at 80°C for 8 hours to synthesize poly(n-butyl acrylate) having isocyanato groups at both ends. The mercapto group-terminated acrylonitrile-methyl methacrylate diblock copolymer (92 parts by weight) synthesized in Produce Example 49 2 was added thereinto, and stirring was performed at 80°C for 16 hours. 14 NMR and GPC analysis confirmed that an (acrylonitrile/methyl methacrylate)-in-butyl acrylate-(acrylonitrile/methyl methacrylate) triblock copolymer was produced (Moy = 122 100, Mn = 78 900, and Mw/Mn = 1.55).

(Example 20: Synthesis of (acrylonitrile/methyl methacrylate)-(n-butyl acrylate/2-methoxyethyl acrylate)-(acrylonitrile/methyl methyl methacrylate) triblock copolymer)

[0138] In a nitrogen atmosphere, dehydrated foluene (500 parts by weight), isophorone diisocyanate (7.22 parts by weight), and dibutyltin bis(isoocyt) thioglycolate) (0.05 parts by weight) were added to the n-butyl acrylate2-methox-yethyl acrylate random copolymer having isocyanate groups at both ends (100 parts by weight) synthesized in Production Example 5, and the mixture was sirred at 80°C for 10 hours to synthesize an n-butyl acrylate12-methoxyethyl acrylate random copolymer having isocyanate groups at both ends. The mercapic group-terminated acrylontificinethyl meth-acrylate random copolymer (474 parts by weight) synthesized in Production Example 2 was added thereinto, and string was performed at 80°C for 15 hours. It MNR and QPC analysis confirmed that an (acrylontificinethyl) meth-

acrylate)-(n-butyl acrylate/2-methoxyethyl acrylate)-(acrylonitrile/methyl methacrylate) triblock copolymer was produced (Mw = 57,700, Mn = 35,500, and Mw/Mn = 1.63).

(Example 21: Synthesis of (acrylonitrile/methyl methacrylate)-vinyl chloride diblock copolymer)

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[0139] The mercapic group-terminated poly(viny) chloride) (141 parts by weight) synthesized in Production Example 6, tetrahydrofuran (300 parts by weight), and lead dioxide (0.5 parts by weight) synthesized in Production Example 1, tetrahydrofuran (300 parts by weight) synthesized in Production Example 2, and the mixture was stirred at 80°C for 10 hours in an air atmosphere. ¹⁴ NMR analysis confirmed that an (acrylonitrile/methyl methacrylate)-vinyl chloride diblock copolymer in which coupling was performed via disulfide bonds was produced, GPC analysis confirmed that Mw = 68,200. Mn = 46,200, and Mw/Mn = 1.4 Mn.

(Example 22: Synthesis of acrylonitrile-n-butyl acrylate-acrylonitrile triblock copolymer)

[0140] In a nitrogen atmosphere, dehydrated toluene (300 parts by weight), ally isocyanate (0.5 parts by weight), and dibuty(tin bis(isocty) thioglycolate) (0.02 parts by weight) were added to the poly(n-butyl acrylate) having mercapto groups at both ends (100 parts by weight) synthesized in Production Example 4, and the mixture was stirred at 80°C for 5 hours. Dibutylith bis(isocoty) thioglycolate) and excess ally isocyanate were removed through a silica gel column (50 parts by weight), and toluene was removed by distillation. Thereby, poly(n-butyl acrylate) having ally groups at both ends via thiourethane bonds was produced. NMR analysis confirmed that the Introduction rate of allyl groups was 89% on the both-ends basis.

[0141] On the other hand, the polyacrylonitrile (100 parts by weight) synthesized in Production Exemple 7 was dissolved in dimethylformamide (300 parts by weight), and 3-isocyanalopropyldimethylsilane (1.4 parts by weight) and dibutylith bisacetylacetonate (0.01 parts by weight) were added thereinto, followed by stirring at 80°C for 6 hours. The reaction solution was concentrated and reprecipitated with methanol. Polyacrylonitrile having a hydrosilyl group at one end was thereby produced. NMR analysis confirmed that the introduction rate of hydrosilyl groups was 88% on the sincle-end basils.

19142] Dimethylformamide (300 parts by weight) was added to a mixture of the resultant polyacrylonifrile having the hydrosilyl group at one end (100 parts by weight) and the poly/n-bulyl acryleta having the abily group at one end, and a 3% by weight xylene solution of platinum-1,3-diviny/tetramethyldisiloxane complex (0.3 parts by weight) was added thereinto. After the mixture was metured at 80° Ct of a hours, dimethylformamide was removed by distillation. Tolluene (200 parts by weight) was added thereto, and insolubles were removed. The solution was poured into methanol (500 parts by weight), and reprecipitation was performed. Thereby, an acrylonitrile-n-butyl acrylate-acrylonitrile triblock co-polymer was produced (five 3 ±28,800, Mn = 73,200, and Mw/Mn = 1.78).

(Comparative Production Example 1: Synthesis of acrylonitrile/n-butyl acrylate random copolymer)

[0143] Into a 2 L reactor equipped with an agitator, a thermometer, a nitrogen gas inlet tube, a dropping funnel, and a reflux condenser, was placed 600 g of distilled water. 0,71 g of sodium dodecyl suifate as an emulsifier, and as monomers, 70 g of acrylonitrile and 30 g of n-butyl acrylate, and the reactor was nitrogen-purged while the reaction mixture was being stifred at 60°C. Next, 1.6 g of 4-fazobsigf-cyanovalent acid) as a polymertzation initiator together with 30 g of distilled water was added into the reactor, and a reaction was initiated. Stirring was performed at 80°C for 10 hours, and an acrylonitrile/h-butyl acrylate random copolymer was produced with a conversion rate of monomers of 89%, GPC measurement confirmed that Mw = 472,000, Mm = 214,000, and Mw/Mn = 2.400.

(Example 23: Polycarbonate thermoplastic resin composition)

[0144] A polycarbonate resin LEXAN 141R-111 (manufactured by GE Plastics Japan, Ltd.) (100 parts by weight) as a thermoplastic resin, IRGANOX HP2215 (manufactured by Ciba Specially Chemicals) (0.5 parts by weight) as a stabilizer, and the block copolymer synthesized in Example 1 (6 parts by weight) were compounded. The resultant composition was exfrusion-kneaded at 280°C with a twin-screw extruder (32 mm, LID = 25.5) and pelletized. The resultant pellets were dired at 80°C for 15 noves, and nijection molding was then performed at 280°C to form a molded object (1/4 inch thick) for evaluating physical properties. The Izod impact strength at 0°C and flame relardancy of the resultant predicted object were evaluated. The results thereof are shown in Table 1 below.

[0145] Additionally, Izod impact strength was measured according to ASTM D256-56, using V-notched specimens, and average values measured at n = 5 were adopted. Flame retardancy was measured according to UL-94 standard.

(Examples 24 to 44)

[0146] Molded objects were formed as in Example 23 except that the block copolymers synthesized in Examples 2 to 22 were used instead of the block copolymer synthesized in Example 1. The Izod impact strength at 0°C and flame retardancy were evaluated. The results thereof are shown in Table 1.

(Comparative Example 1)

[0147] A molded object was formed as in Example 23 except that the random copolymer synthesized in Comparative Production Example 1 was used instead of the block copolymer synthesized in Example 1. The Izod impact strength at 0°C and flame retardancy were evaluated. The results thereof are shown in Table 1.

(Comparative Example 2)

i [0148] A molded object was formed as in Example 23 without compounding the block copolymer synthesized in Example 1. The Izod impact strength at 0°C and flame retardancy were evaluated. The results thereof are shown in Table 1.

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Example	Composition	Izod impact strength (kJ/m²)	Flame retardancy
Example 23	Example 1	10.5	V-2
Example 24	Example 2	9.7	V-2
Example 25	Example 3	11.4	V-2
Example 26	Example 4	11.5	V-1
Example 27	Example 5	15.2	V-1
Example 28	Example 6	16.1	V-1
Example 29	Example 7	11.8 .	V-1
Example 30	Example 8	11.8	V-1
Example 31	Example 9	14.9	V-1
Example 32	Example 10	12.5	V-1
Example 33	Example 11	10.1	V-0
Example 34	Example 12	10.0	V-2
Example 35	Example 13	14.7	V-2
Example 36	Example 14	9,1	V-2
Example 37	Example 15	13.3	V-2
Example 38	Example 16	12.0	V-1
Example 39	Example 17	7.2	V-0
Example 40	Example 18	8.8	V-2
Example 41	Example 19	17.1	V-1
Example 42	Example 20	16.1	V-1
Example 43	Example 21	7.7	V-0
Example 44	Example 22	10.2	V-1
Comparative Example 1	Comparative Production Example 1	4.1	netV
Comparative Example 2	-	2.9	notV

[0149] As is evident from Table 1, the thermoplastic resin compositions in which the block copolymers of the present

invention are compounded are excellent in impact strength at low temperatures and flame retardancy.

(Example 45: Polyester thermoplastic resin composition)

5 (9150) A poly(butylene terephthalate) resin DURANEX 2002 (manufactured by Polyplastic Co., Ltd.) (80 parts by weight) as a thempolastic resin, Topanol CA (manufactured by Lipre Co., Ltd.) (0.3 parts by weight) as a phenolic antioxidant, Adekasulabu PEP-36 (manufactured by Asahi Denka Co., Ltd.) (0.3 parts by weight) as a HALS, and the block copolymer synthesized in Example 1 (20 parts by weight) were compounded. Using a brim-screw extruct of mr. LD = 25.5), the resultant composition was extrusion-kneaded at 24°C and pelletized. The resultant pellets were dried at 80°C for 15 hours, and injection molding was then performed at 250°C to form a molded object (1/4 inch thick) for evaluating physical properties. The Izod impact strength at 0°C of the resultant molded object was evaluated. The result thereof is shown in Table 2 below.

(Examples 46 to 66)

[0151] Molded objects were formed as in Example 45 except that the block copolymers synthesized in Examples 2 to 22 were used instead of the block copolymer synthesized in Example 1. The tzod impact strength at 0°C of each resultant molded object was evaluated. The results thereof are shown in

20 Table 2.

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(Comparative Example 3)

[0152] A moided object was formed as in Example 45 except that the random copolymer synthesized in Comparative 25 Production Example 1 was used instead of the block copolymer synthesized in Example 1. The Izod impact strength at 0°C was evaluated. The result thereof is shown in Table 2.

(Comparative Example 4)

[0153] A molided object was formed as in Example 45 without compounding the block copolymer synthesized in Example 1. The Izod impact strength at 0°C was evaluated. The result thereof is shown in Table 2.

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Example	Composition	Izod impact strength (kJ/m²)
Example 45	Example 1	9.6
Example 46	Example 2	8.5
Example 47	Example 3	9.3
Example 48	Example 4	10.1
Example 49	Example 5	11.5
Example 50	Example 6	11.8
Example 51	Example 7	10.1
Example 52	Example 8	9.1
Example 53	Example 9	12.5
Example 54	Example 10	10.9
Example 55	Example 11	9.1
Example 56	Example 12	8.3
Example 57	Example 13	11.1
Example 58	Example 14	9.0
Example 59	Example 15	10.8
Example 60	Example 16	10.7

(Continued)

Example	Composition	Izod impact strength (kJ/m²)
Example 61	Example 17	8,1
Example 62	Example 18	8.3
Example 63	Example 19	13.2
Example 64	Example 20	13.7
Example 65	Example 21	7.8
Example 66	Example 22	9.3
Comparative Example 3	Comparative Production Example 1	3.1
Comparative Example 4	-	2.8

[0154] As is evident from Table 2, the thermoplastic resin compositions in which the block copolymers of the present invention are compounded are excellent in impact strength at low temperatures.

(Example 67: Elastomer composition)

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[0155] An acrylic rubber AR42W (manufactured by ZEON Corporation) (100 parts by weight) and the block copolymer synthesized in Example 1 (100 parts by weight) were compounded and melt-kneaded with a Laboplastomiil at 190°C. After 3 minutes, as a crosslinking agont, ammonium benzoate (3 parts by weight) was added thereinto, and melt-kneading was further performed with the Laboplastomiil at 190°C. The resultant sample block was thermopress-molded at 190°C, and molded objects with a thickness of 2 mm for evaluating physical properties were formed. The resultant molded objects were crosslinked by heating at 150°C for 2 hours. With respect to the molded objects, tensile strength at break and elongation at break at 0°C, and oil resistance were evaluated. The results thereof are shown in Table 3 hellow.

[0156] Additionally, tensile strength at break and elongation at break were measured according to JIS K6251, at 0°C. With respect to oil resistance, according to JIS C232, molded objects were immersed in transformer oil B at 70°C for 4 hours, and the oil resistance was measured based on the rates of change in weight.

(Examples 68 to 88)

[0157] Molded objects were formed as in Example 67 except that the block copolymers synthesized in Examples 2 to 22 were used instead of the block copolymers synthesized in Example 1. The tensile strength at break and elongation at break at 07°C, and oil resistance were evaluated. The results thereof are shown in Table 3.

(Comparative Example 5)

[0158] Molded objects were formed as in Example 67 except that the random copolymer synthesized in Comparative Production Example 1 was used instead of the block copolymer synthesized in Example 1. The tensile strength at break and elongation at break of °C, and oil resistance were evaluated. The results thereof are shown in Table 3.

45 (Comparative Example 6)

[0159] Molded objects were formed as in Example 67 without compounding the block copolymer synthesized in Example 1. The tensite strength at break and elongation at break at 0°C, and oil resistance were evaluated. The results thereof are shown in Table 3.

[Table 3]

Example	Composition	Tensile strength at break (MPa)	Elongation at break (%)	Oil resistance (%)
Example 67	Example 1	4.1	340	11
Example 68	Example 2	5.2	310	9

(Table 3) (continued)

Example	Composition	Tensile strength at break (MPa)	Elongation at break (%)	Oil resistance (%)
Example 69	Example 3	4.8	300	7
Example 70	Example 4	3.5	420	4
Example 71	Example 5	7.1	220	12
Example 72	Example 6	6.6	230	11
Example 73	Example 7	5.9	280	10
Example 74	Example 8	4.5	270	10
Example 75	Example 9	6.0	240	6
Example 76	Example 10	6.9	250	7
Example 77	Example 11	8.1	200	4
Example 78	Example 12	4.0	300	13
Example 79	Example 13	5.0	270	14
Example 80	Example 14	3.7	360	14
Example 81	Example 15	4.2	350	13
Example 82	Example 16	4.5	330	11
Example 83	Example 17	3.0	470	8
Example 84	Example 18	3.9	370	15
Example 85	Example 19	4.0	370	14
Example 86	Example 20	6.2	210	16
Example 87	Example 21	3.3	440	8
Example 88	Example 22	3.7	420	11
Comparative Example 5	Comparative Production Example 1	1.4	500	28
Comparative Example 6	-	1.6	490	21

[0160] As is evident from Table 3, the elastomer compositions in which the block copolymers of the present invention are compounded are excellent in strength at low temperatures and oil resistance.

Industrial Applicability

[0161] As described above, in accordance with the present invention, it is possible to produce (meth)acrylonitrile-based block copolymers which are excellent in heat resistance, weatherability, oil resistance, flame retardancy, and low-temperature resistance. The block copolymers can be prepared by water-based polymerization and the purification step can be simplified, and thus economical production is enabled. The thermoplastic resin compositions and elastomer compositions containing the block copolymers can be used in various applications, such as films, sheets, tapes, hoses, tubes, gaskets, packings, grips, containers, various molded objects, sealants, damping materials, pressure-sensitive adhesives, coating materials, potting materials, and textiles.

Claims

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 A block copolymer comprising a polymer block (A) and a polymer block (B), the block copolymer being produced by forming the polymer block (A) by reversible addition-fragmentation chain transfer polymerization in the presence of a thiocarbonylthio group-containing compound, and then by forming the polymer block (B),

wherein the polymer block (A) is prepared by (co)polymerizing 50% to 100% by weight of at least one monomer selected from the group consisting of acryloritrile and methacrylonitrile and 50% to 0% by weight of at least one monomer selected from the group consisting of methacrylate seters, styrene, and α-methylstyrene, and the polymer block (B) is prepared by (co)polymerizing at least one monomer selected from the group consisting of acrylic acid, acrylate esters, winyl acetate, styrene, α-methylstyrene, buta-diene, isoprane, and vinyl chioride.

- A block copolymer produced by separately forming the polymer block (A) and the polymer block (B) described
 above, each by reversible addition-fragmentalian chain transfer polymerization in the presence of a thiocarbonylthio group-containing compound, and then by opuling the polymer block (A) and the polymer block (B).
 - 3. A block copolymer prepared by coupling of the block copolymer according to Claim 1.
- 4. The block copolymer according to any one of Cleims 1 to 3, wherein the thiocarbonyithio group-containing compound is at least one compound selected from the group consisting of a compound represented by general formula (1):

$$\left(\begin{array}{c}
S\\Z^1-C-S\\p\end{array}\right)_{p}R^1 \qquad (1)$$

(wherein R¹ is a p-valent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, phosphorus, and metal atoms, or which may be a polymer; Z¹ is a hydrogen atom, natom or monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, and phosphorus atoms, or which may be a polymer; when plural Z¹s are present, the plural Z¹s are present, t

$$Z^{2} \begin{pmatrix} S \\ C - S - R^{2} \end{pmatrix}_{q}$$
 (2)

- (wherein R² is a monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, phosphorus, and metal atoms, or which may be a polymer, Z² is an oxygen atom (when q = 2), sulfur atom (when q = 2), nitrogen atom (when q = 3), or q-valent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, and phosphorus atoms, or which may be a polymer; plural R²s may be the same or different; and q is an integer of 2 or more).
 - The block copolymer according to any one of Claims 1 to 4, wherein the thiocarbonylthio group-containing compound is a compound represented by general formula (3):

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$$S$$
 Z^1 — C — S — R^2 (3)

(wherein \mathbb{R}^2 is a monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, phosphorus, and metal atoms, or which may be a polymer, and \mathbb{Z}^1 is a hydrogen atom, halogen atom, or monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, silicon, and phosphorus atoms, or which may be a polymer).

6. The block copolymer according to Claim 2, wherein the polymerization reaction for forming the polymer block (A) is performed in the presence of the thiocarbonylithic group-containing compound represented by general formula (3) described above, and the polymerization reaction for forming the polymer block (B) is performed in the presence of a thiocarbonylithic group-containing compound represented by general formula (4):

$$S$$
 S Z^{1} — $C-S-R^{3}-S-C-Z^{1}$ S

(wherein R³ is a divalent organic group of 1 or more carbon atoms which may contain one of nitrogen, oxygen, sulfur, halogen, sillicon, phosphorus, and metal atoms, or which may be a polymer; each Z¹ is a hydrogen atom, halogen atom, or monovalent organic group of 1 or more carbon atoms which may contain one of nitrogen. oxygen, sulfur, halogen, sillicon, and phosphorus atoms, or which may be a polymer; and two Z¹s may be the same or different).

- The block copolymer according to either Claim 2 or 6, wherein the molecular weight distribution of the polymer block (B) is 1.8 or less, the molecular weight distribution being determined by gel permeation chromatography analysis.
- The block copolymer according to any one of Claims 2, 3, 6, and 7, wherein the thiocarbonythio group of the block copolymer is converted into a mercapto group or a mercaptide group, and said coupling is performed by a reaction of the mercapto coroup or the mercaptide group.
- 9. The block copolymer according to Claim 8, wherein the thiocarbonyithio group of the thiocarbonyithio group-containing block copolymer is converted into the mercapto group or the mercaptide group by a reaction with a processing agent comprising at least one compound selected from the group consisting of bases, acids, and hydrogen-through bond-containing compounds.
- 48 10. The block copolymer according to Ctalm 9, wherein the processing agent is at least one compound selected from the group consisting of ammonia, primary amine compounds with a boiling point of 100°C or less, secondary amine compounds with a boiling point of 100°C or less, and hindered amine light stabilizers (HALSs).
- The block copolymer according to any one of Claims 8 to 10, wherein said coupling is performed by the formation of disulfide bonds using an oxidizing agent.
 - 12. The block copolymer according to any one of Claims 8 to 10, wherein said coupling is performed using a compound having at least two groups in each molecule, each group reactive with the mercapto group or the mercaptide group to form a bond.
 - 13. The block copolymer according to Claim 12, wherein said coupling is performed using a compound having at least two isocyanato groups in each molecule.

- 14. The block copolymer according to any one of Claims 8 to 10, wherein a functional group is introduced into the mercapto group or the mercaptide group for performing said coupling.
- 15. The block copolymer according to Claim 14, wherein said coupling is performed using at least one functional group selected from the group consisting of a crosslinkable sityl group, unsaturated groups, and a hydroxyl group.
 - 16. The block copolymer according to any one of Claims 1 to 15, wherein the polymer block (A) is prepared by polymerizing 80% to 100% by weight of at least one monomer selected from the group consisting of acrylonitrile and methacrylonitrile and 20% to 0% by weight of at least one monomer selected from the group consisting of methacrylonite esters, styrene, and α-methylstyrene.

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- 17. The block copolymer according to Claim 16, wherein the polymer block (A) is prepared by polymerizing 100% by weight of at least one monomer selected from the group consisting of acrylonitrile and methacrylonitrile.
- 18. The block copolymer according to any one of Claims 1 to 17, wherein the polymer block (B) is prepared by polymerizing an acrylate ester.
 - 19. The block copolymer according to any one of Claims 1 to 18, wherein the molecular weight distribution of the polymer block (A) is 1.8 or less, the molecular weight distribution being determined by get permeation chromatography analysis.
 - 20. The block copolymer according to any one of Claims 1 to 19, wherein the molecular weight distribution of the block copolymer comprising the polymer block (A) and the polymer block (B) is 1.8 or less, the molecular weight distribution being determined by eql permeation chromatography analysis.
 - The block copolymer according to any one of Claims 1 to 20, wherein the glass transition temperature of the polymer block (A) is 50°C or more.
- 22. The block copolymer according to any one of Claims 1 to 21, wherein the glass transition temperature of the polymer block (B) is 30°C or less.
 - A thermoplastic resin composition comprising a thermoplastic resin and the block copolymer according to any one
 of Claims 1 to 22.
- 35 24. An elastomer composition comprising a synthetic rubber and the block copolymer according to any one of Claims 1 to 22

INTERNATIONAL SEARCH REPORT

Intertracional application No.
PCT/JP02/06656

A. CLASSIFICATION OF SUBJECT MATTER Int.Cl ¹ C09F293/00		-		
According to International Patent Classification (IPC) or to both national classification and IPC				
B. FIELDS SEARCHED				
Minimum documentation searched (classification system followed Int.Cl ² C08F293/00	by classification symbols)			
Documentation searched other than minimum documentation to the	e extent that such documents are included in the	e fields searched		
Electronic data base consulted during the international search (name	se of data base and, where practicable, search te	rris used)		
C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category* Citation of document, with indication, where as	propriate, of the relevant passages Re	levant to claim No.		
F,A JF 2002-241568 A (Raneka Cor 28 August, 2002 (28.08.02), Claims	p.),	1-24		
A JP 2000-154329 A (Kaneka Co: 06 June, 2000 (06.06.00), Claims (Family: none)		1-24		
Further documents are listed in the continuation of Box C.	See patent family annex.			
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